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(54) Title: MANIPULATION OF FLAVONOID BIOSYNTHESIS IN PLANTS

(57) Abstract: The present invention relates to nucleic acids and nucleic acid fragments encoding amino acid sequences for flavonoid biosynthetic enzymes in plants, and the use thereof for the modification of flavonoid biosynthesis in plants. More particularly, the flavonoid biosynthetic enzyme is selected from the group consisting of chalcone isomerase (CHI), chalcone synthase (CHS), chalcone reductase (CHR), dihydroflavonol 4-reductase (DFR), leucoanthocyanidin reductase (LCR), flavonoid 3', 5' hydrolase (F3'5'H), flavanone 3-hydroxylase (F3H), flavonoid 3'-hydroxylase (F3'H), phenylalanine ammonia-lyase (PAL) and vestitone reductase (VR), and functionally active fragments and variants thereof.



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## MANIPULATION OF FLAVONOID BIOSYNTHESIS IN PLANTS

The present invention relates to nucleic acids and nucleic acid fragments encoding amino acid sequences for flavonoid biosynthetic enzymes in plants, and the use thereof for the modification of flavonoid biosynthesis in plants.

5        Flavonoids constitute a relatively diverse family of aromatic molecules that are derived from phenylalanine and malonyl-coenzyme A (CoA, via the fatty acid pathway). These compounds include six major subgroups that are found in most higher plants: the chalcones, flavones, flavonols, flavandiols, anthocyanins and condensed tannins (or proanthocyanidins). A seventh group, the aurones, is  
10        widespread, but not ubiquitous.

      Some plant species also synthesize specialised forms of flavonoids, such as the isoflavonoids that are found in legumes and a small number of non-legume plants. Similarly, sorghum, maize and gloxinia are among the few species known to synthesize 3-deoxyanthocyanins (or phlobaphenes in the polymerised form).  
15        The stilbenes which are closely related to flavonoids, are synthesised by another group of unrelated species that includes grape, peanut and pine.

      Besides providing pigmentation to flowers, fruits, seeds, and leaves, flavonoids also have key roles in signalling between plants and microbes, in male fertility of some species, in defense as antimicrobial agents and feeding  
20        deterrents, and in UV protection.

      Flavonoids also have significant activities when ingested by animals, and there is great interest in their potential health benefits, particularly for compounds such as isoflavonoids, which have been linked to anticancer benefits, and stilbenes that are believed to contribute to reduced heart disease.

25        The major branch pathways of flavonoid biosynthesis start with general phenylpropanoid metabolism and lead to the nine major subgroups: the colorless chalcones, aurones, isoflavonoids, flavones, flavonols, flavandiols, anthocyanins, condensed tannins, and phlobaphene pigments. The enzyme phenylalanine



ammonia-lyase (PAL) of the general phenylpropanoid pathway will lead to the production of cinnamic acid. Cinnamate-4-hydroxylase (C4H) will produce p-coumaric acid which will be converted through the action of 4-coumaroyl:CoA-ligase (4CL) to the production of 4-coumaroyl-CoA and malonyl-CoA. The first  
5 committed step in flavonoid biosynthesis is catalyzed by chalcone synthase (CHS), which uses malonyl CoA and 4-coumaroyl CoA as substrates. Chalcone reductase (CHR) balances the production of 5-hydroxy- or 5-deoxyflavonoids. The next enzyme, chalcone isomerase (CHI) catalyses ring closure to form a flavanone, but the reaction can also occur spontaneously. Other enzymes in the  
10 pathway are: flavanone 3-hydroxylase (F3H), dihydroflavonol 4-reductase (DFR), flavonoid 3'-hydroxylase (F3'H) and flavonoid 3', 5' hydroxylase (F3'5'H).

The *Arabidopsis* *BANYULS* gene encodes a DFR-like protein that may be a leucoanthocyanidin reductase (LCR) that catalyzes an early step in condensed tannin biosynthesis. Condensed tannins are plant polyphenols with protein-  
15 precipitating and antioxidant properties, synthesized by the flavonoid pathway. Their chemical properties include protein binding, metal chelation, anti-oxidation, and UV-light absorption. As a result condensed tannins inhibit viruses, microorganisms, insects, fungal pathogens, and monogastric digestion. Moderate amounts of tannins improve forage quality by disrupting protein foam and  
20 conferring protection from rumen pasture bloat. Bloat is a digestive disorder that occurs on some highly nutritious forage legumes such as alfalfa (*Medicago sativa*) and white clover (*Trifolium repens*). Moderate amounts of tannin can also reduce digestion rates in the rumen and can reduce parasitic load sufficiently to increase the titre of amino acids and small peptides in the small intestine without  
25 compromising total digestion.

Vestitone reductase (VR) is the penultimate enzyme in medicarpin biosynthesis. Medicarpin, a phytoalexin, has been associated with plant resistance to fungal pathogens.

While nucleic acid sequences encoding some flavonoid biosynthetic  
30 enzymes CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and VR have been isolated for certain species of plants, there remains a need for materials useful in

modifying flavonoid biosynthesis; in modifying protein binding, metal chelation, anti-oxidation, and UV-light absorption; in modifying plant pigment production; in modifying plant defense to biotic stresses such as viruses, micro-organisms, insects or fungal pathogens; in modifying forage quality, for example by disrupting  
5 protein foam and/or conferring protection from rumen pasture bloat, particularly in forage legumes and grasses, including alfalfa, medics, clovers, ryegrasses and fescues, and for methods for their use.

It is an object of the present invention to overcome, or at least alleviate, one or more of the difficulties or deficiencies associated with the prior art.

10 In one aspect, the present invention provides substantially purified or isolated nucleic acids or nucleic acid fragments encoding the flavonoid biosynthetic enzymes CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and VR from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species and functionally active fragments and variants thereof.

15 The present invention also provides substantially purified or isolated nucleic acids or nucleic acid fragments encoding amino acid sequences for a class of proteins which are related to CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and VR and functionally active fragments and variants thereof. Such proteins are referred to herein as CHI-like, CHS-like, CHR-like, DFR-like, LCR-like, F3'5'H-like,  
20 F3H-like, F3'H-like, PAL-like and VR-like, respectively.

The individual or simultaneous enhancement or otherwise manipulation of CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and/or VR or like gene activities in plants may enhance or otherwise alter flavonoid biosynthesis; may enhance or otherwise alter the plant capacity for protein binding, metal chelation,  
25 anti-oxidation or UV-light absorption; may enhance or reduce or otherwise alter plant pigment production; may modify plant defense to biotic stresses such as viruses, micro-organisms, insects or fungal pathogens; and/or may modify forage quality, for example by disrupting protein foam and/or conferring protection from rumen pasture bloat.

The individual or simultaneous enhancement or otherwise manipulation of CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and/or VR or like gene activities in plants has significant consequences for a range of applications in, for example, plant production and plant protection. For example, it has applications in

5 increasing plant tolerance and plant defense to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; in improving plant forage quality, for example by disrupting protein foam and in conferring protection from rumen pasture bloat; in reducing digestion rates in the rumen and reducing parasitic load; in the production of plant compounds leading to health benefits, such as

10 isoflavonoids, which have been linked to anticancer benefits, and stilbenes that are believed to contribute to reduced heart disease.

Methods for the manipulation of CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and/or VR or like gene activities in plants, including legumes such as clovers (*Trifolium* species), lucerne (*Medicago sativa*) and grass species such as

15 ryegrasses (*Lolium* species) and fescues (*Festuca* species) may facilitate the production of, for example, forage legumes and forage grasses and other crops with enhanced tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; altered pigmentation in flowers; forage legumes with enhanced herbage quality and bloat-safety; crops with enhanced isoflavonoid

20 content leading to health benefits.

The clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species may be of any suitable type, including white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*), alfalfa (*Medicago sativa*), Italian or annual ryegrass (*Lolium multiflorum*), perennial ryegrass (*Lolium perenne*), tall fescue (*Festuca arundinacea*), meadow fescue (*Festuca pratensis*) and red fescue (*Festuca rubra*). Preferably the species is a clover or a ryegrass, more preferably white clover (*T. repens*) or perennial ryegrass (*L. perenne*). White clover (*Trifolium repens* L.) and perennial ryegrass (*Lolium perenne* L.) are key pasture legumes and grasses,

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30 respectively, in temperate climates throughout the world. Perennial ryegrass is also an important turf grass.

The nucleic acid or nucleic acid fragment may be of any suitable type and includes DNA (such as cDNA or genomic DNA) and RNA (such as mRNA) that is single- or double-stranded, optionally containing synthetic, non-natural or altered nucleotide bases, and combinations thereof.

5       The term “isolated” means that the material is removed from its original environment (eg. the natural environment if it is naturally occurring). For example, a naturally occurring nucleic acid present in a living plant is not isolated, but the same nucleic acid separated from some or all of the coexisting materials in the natural system, is isolated. Such nucleic acids could be part of a vector and/or  
10 such nucleic acids could be part of a composition, and still be isolated in that such a vector or composition is not part of its natural environment.

Such nucleic acids or nucleic acid fragments could be assembled to form a consensus contig. As used herein, the term “consensus contig” refers to a nucleotide sequence that is assembled from two or more constituent nucleotide  
15 sequences that share common or overlapping regions of sequence homology. For example, the nucleotide sequence of two or more nucleic acids or nucleic acid fragments can be compared and aligned in order to identify common or overlapping sequences. Where common or overlapping sequences exist between two or more nucleic acids or nucleic acid fragments, the sequences (and thus their  
20 corresponding nucleic acids or nucleic acid fragments) can be assembled into a single contiguous nucleotide sequence.

In a preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a CHI or CHI-like protein includes a nucleotide sequence selected from the group consisting of  
25 (a) sequences shown in Figures 1, 3, 4, 6, 7, 9, 10, 12, 122 and 127 hereto (Sequence ID Nos: 1, 3 to 7, 8, 10 to 12, 13, 15 and 16, 17, 19 to 22, 307, and 309, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a CHS or CHS-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 13, 15, 16, 18, 19, 21, 22, 24, 25, 27, 28, 30, 31, 33, 34, 137, 142, 147, 152, 157 and 162 hereto (Sequence ID Nos: 23, 25 to 63, 64, 66 to 68, 69, 71 to 77, 78, 80 to 90, 91, 93 and 94, 95, 97 to 100, 101, 103 to 105, 106, 313, 315, 317, 319, 321, and 323, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a CHR or CHR-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 36, 38, 40, 41, 43 and 132 hereto (Sequence ID Nos: 108, 110, 112 to 116, 117, 119 to 134, and 311, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a DFR or DFR-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 44, 46, 47, 49, 50, 52, 54, 55, 57, 59, 61, 62, 64, 101, 103, 104, 106, 117 and 167 hereto (Sequence ID Nos: 135, 137 to 146, 147, 149 to 152, 153, 155, 157 and 158, 159, 161, 163, 165 to 167, 168, 170 to 184, 286, 288 to 292, 293, 295 to 297, 305, and 325, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an LCR or LCR-like protein includes a nucleotide sequence selected from the group

consisting of (a) sequences shown in Figures 65 and 67 hereto (Sequence ID Nos: 185 and 187 to 193, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an F3'5'H or F3'5'H-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 68, 70 and 72 hereto (Sequence ID Nos: 194, 196, and 198 to 201, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an F3H or F3H-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 73, 75, 76, 78, 107, 109, 111 and 172 hereto (Sequence ID Nos: 202, 204 to 244, 245, 247, 298, 300 to 302, 303, and 327, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an F3'H or F3'H-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 80 and 82 hereto (Sequence ID Nos: 249, and 251 and 252, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an PAL or PAL-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 83, 85, 86, 88, 89, 91, 93, 95, 97,  
5 177, 182 and 187 hereto (Sequence ID Nos: 253, 255 to 257, 258, 260 to 267, 268, 270, 272, 274, 276 and 277, 329, 331, and 333, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

10 In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding an VR or VR-like protein includes a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 98, 100 and 192 hereto (Sequence  
ID Nos: 278, 280 to 285, and 335, respectively); (b) complements of the  
15 sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

By "functionally active" in relation to nucleic acids it is meant that the fragment or variant (such as an analogue, derivative or mutant) encodes a  
20 polypeptide which is capable of modifying flavonoid biosynthesis in a plant. Such variants include naturally occurring allelic variants and non-naturally occurring variants. Additions, deletions, substitutions and derivatizations of one or more of the nucleotides are contemplated so long as the modifications do not result in loss of functional activity of the fragment or variant. Preferably the functionally active  
25 fragment or variant has at least approximately 80% identity to the relevant part of the above mentioned nucleotide sequence, more preferably at least approximately 90% identity, even more preferably at least approximately 95% identity, most preferably at least approximately 98% homology. Such functionally active variants and fragments include, for example, those having nucleic acid changes which  
30 result in conservative amino acid substitutions of one or more residues in the corresponding amino acid sequence. Preferably the fragment has a size of at least

10 nucleotides, more preferably at least 15 nucleotides, most preferably at least 20 nucleotides.

Nucleic acids or nucleic acid fragments encoding at least a portion of several CHI, CHS, CHR, DFR, LCR, F3'5'H, F3H, F3'H, PAL and VR have been  
5 isolated and identified. The nucleic acids or nucleic acid fragments of the present invention may be used to isolate cDNAs and genes encoding homologous proteins from the same or other plant species. Isolation of homologous genes using sequence-dependent protocols, such as methods of nucleic acid hybridisation, and methods of DNA and RNA amplification as exemplified by various uses of nucleic  
10 acid amplification technologies (e.g. polymerase chain reaction, ligase chain reaction), is well known in the art.

For example, genes encoding other CHI or CHI-like, CHS or CHS-like, CHR or CHR-like, DFR or DFR-like, LCR or LCR-like, F3'5'H or F3'5'H-like, F3H or F3H-like, F3'H or F3'H-like, PAL or PAL-like and VR or VR-like proteins, either as  
15 cDNAs or genomic DNAs, may be isolated directly by using all or a portion of the nucleic acids or nucleic acid fragments of the present invention as hybridisation probes to screen libraries from the desired plant employing the methodology well known to those skilled in the art. Specific oligonucleotide probes based upon the nucleic acid sequences of the present invention may be designed and synthesized  
20 by methods known in the art. Moreover, the entire sequences may be used directly to synthesize DNA probes by methods known to the skilled artisan such as random primer DNA labelling, nick translation, or end-labelling techniques, or RNA probes using available *in vitro* transcription systems. In addition, specific primers may be designed and used to amplify a part or all of the sequences of the present  
25 invention. The resulting amplification products may be labelled directly during amplification reactions or labelled after amplification reactions, and used as probes to isolate full-length cDNA or genomic fragments under conditions of appropriate stringency.

In addition, short segments of the nucleic acids or nucleic acid fragments of  
30 the present invention may be used in amplification protocols to amplify longer nucleic acids or nucleic acid fragments encoding homologous genes from DNA or



RNA. For example, polymerase chain reaction may be performed on a library of cloned nucleic acid fragments wherein the sequence of one primer is derived from the nucleic acid sequences of the present invention, and the sequence of the other primer takes advantage of the presence of the polyadenylic acid tracts to the 3' end of the mRNA precursor encoding plant genes. Alternatively, the second primer sequence may be based upon sequences derived from the cloning vector. For example, those skilled in the art can follow the RACE protocol [Frohman *et al.* (1988) *Proc. Natl. Acad. Sci. USA* 85:8998, the entire disclosure of which is incorporated herein by reference] to generate cDNAs by using PCR to amplify copies of the region between a single point in the transcript and the 3' or 5' end. Using commercially available 3' RACE and 5' RACE systems (BRL), specific 3' or 5' cDNA fragments may be isolated [Ohara *et al.* (1989) *Proc. Natl. Acad. Sci. USA* 86:5673; Loh *et al.* (1989) *Science* 243:217, the entire disclosures of which are incorporated herein by reference]. Products generated by the 3' and 5' RACE procedures may be combined to generate full-length cDNAs.

In a second aspect of the present invention there is provided a substantially purified or isolated polypeptide from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species, selected from the group consisting of CHI and CHI-like, CHS and CHS-like, CHR and CHR-like, DFR and DFR-like, LCR and LCR-like, F3'5'H and F3'5'H-like, F3H and F3H-like, F3'H and F3'H-like, PAL and PAL-like, VR and VR-like; and functionally active fragments and variants thereof.

The clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species may be of any suitable type, including white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*), alfalfa (*Medicago sativa*), Italian or annual ryegrass (*Lolium multiflorum*), perennial ryegrass (*Lolium perenne*), tall fescue (*Festuca arundinacea*), meadow fescue (*Festuca pratensis*) and red fescue (*Festuca rubra*). Preferably the species is a clover or a ryegrass, more preferably white clover (*T. repens*) or perennial ryegrass (*L. perenne*).

In a preferred embodiment of this aspect of the invention, the substantially purified or isolated CHI or CHI-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown in Figures 2, 5, 8, 11, 123 and 128 hereto (Sequence ID Nos: 2, 9, 14, 18, 308, and 310, respectively),  
5 and functionally active fragments and variants thereof.

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated CHS or CHS-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown in Figures 14, 17, 20, 23, 26, 29, 32, 35, 138, 143, 148, 153, 158 and 163 hereto  
10 (Sequence ID Nos: 24, 65, 70, 79, 92, 96, 102, 107, 314, 316, 318, 320, 322, and 324, respectively), and functionally active fragments and variants thereof.

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated CHR or CHR-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown in  
15 Figures 37, 39, 42 and 133 hereto (Sequence ID Nos: 109, 111, 118, and 312, respectively), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated DFR or DFR-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown in  
20 Figures 45, 48, 51, 53, 56, 58, 60, 63, 102, 105, 118 and 168 hereto (Sequence ID Nos: 136, 148, 54, 156, 160, 162, 164, 169, 287, 294, 306, and 326, respectively), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated LCR or LCR-like polypeptide includes an amino  
25 acid sequence shown in Figure 66 hereto (Sequence ID No: 186), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated F3'5'H or F3'5'H-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown

in Figures 69 and 71 hereto (Sequence ID Nos: 195 and 197, respectively), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated F3H or F3H-like polypeptide includes an amino acid sequence selected from the group consisting of the sequences shown in  
5 Figures 74, 77, 79, 108, 112 and 173 hereto (Sequence ID Nos: 203, 246, 248, 299, 304, and 328, respectively), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the  
10 substantially purified or isolated F3'H or F3'H-like polypeptide includes an amino acid sequence shown in Figure 81 hereto (Sequence ID No: 250), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated PAL or PAL-like polypeptide includes an amino  
15 acid sequence selected from the group consisting of the sequences shown in Figures 84, 87, 90, 92, 94, 96, 178, 183 and 188 hereto (Sequence ID Nos: 254, 259, 269, 271, 273, 275, 330, 332, and 334, respectively), and functionally active fragments and variants thereof.

In a still further preferred embodiment of this aspect of the invention, the  
20 substantially purified or isolated VR or VR-like polypeptide includes an amino acid sequence shown in Figures 99 and 193 hereto (Sequence ID Nos: 279 and 336, respectively), and functionally active fragments and variants thereof.

By "functionally active" in relation to polypeptides it is meant that the fragment or variant has one or more of the biological properties of the proteins  
25 CHI, CHI-like, CHS, CHS-like, CHR, CHR-like, DFR, DFR-like, LCR, LCR-like, F3'5'H, F3'5'H-like, F3H, F3H-like, F3'H, F3'H-like, PAL, PAL-like, VR and VR-like, respectively. Additions, deletions, substitutions and derivatizations of one or more of the amino acids are contemplated so long as the modifications do not result in loss of functional activity of the fragment or variant. Preferably the functionally

active fragment or variant has at least approximately 60% identity to the relevant part of the above mentioned amino acid sequence, more preferably at least approximately 80% identity, even more preferably at least approximately 90% identity most preferably at least approximately 95% homology. Such functionally  
5 active variants and fragments include, for example, those having conservative amino acid substitutions of one or more residues in the corresponding amino acid sequence. Preferably the fragment has a size of at least 10 amino acids, more preferably at least 15 amino acids, most preferably at least 20 amino acids.

In a further embodiment of this aspect of the invention, there is provided a  
10 polypeptide recombinantly produced from a nucleic acid or nucleic acid fragment according to the present invention. Techniques for recombinantly producing polypeptides are well known to those skilled in the art.

Availability of the nucleotide sequences of the present invention and deduced amino acid sequences facilitates immunological screening of cDNA  
15 expression libraries. Synthetic peptides representing portions of the instant amino acid sequences may be synthesized. These peptides may be used to immunise animals to produce polyclonal or monoclonal antibodies with specificity for peptides and/or proteins including the amino acid sequences. These antibodies may be then used to screen cDNA expression libraries to isolate full-length cDNA  
20 clones of interest.

A genotype is the genetic constitution of an individual or group. Variations in genotype are important in commercial breeding programs, in determining parentage, in diagnostics and fingerprinting, and the like. Genotypes can be readily described in terms of genetic markers. A genetic marker identifies a  
25 specific region or locus in the genome. The more genetic markers, the finer defined is the genotype. A genetic marker becomes particularly useful when it is allelic between organisms because it then may serve to unambiguously identify an individual. Furthermore, a genetic marker becomes particularly useful when it is based on nucleic acid sequence information that can unambiguously establish a  
30 genotype of an individual and when the function encoded by such nucleic acid is known and is associated with a specific trait. Such nucleic acids and/or nucleotide

sequence information including single nucleotide polymorphisms (SNPs), variations in single nucleotides between allelic forms of such nucleotide sequence, may be used as perfect markers or candidate genes for the given trait.

Applicants have identified a number of SNPs of the nucleic acids or nucleic acid fragments of the present invention. These are indicated (marked with grey on the black background) in the figures that show multiple alignments of nucleotide sequences of nucleic acid fragments contributing to consensus contig sequences. See for example, Figures 3, 6, 9, 12, 15, 18, 21, 24, 27, 30, 33, 40, 43, 46, 49, 54, 61, 64, 67, 72, 75, 82, 85, 88, 97, 100, 103, 106 and 109 hereto (Sequence ID Nos: 3 to 7, 10 to 12, 15 and 16, 19 to 22, 25 to 63, 66 to 68, 71 to 77, 80 to 90, 93 and 94, 97 to 100, 103 to 105, 112 to 116, 119 to 134, 137 to 146, 149 to 152, 157 and 158, 165 to 167, 170 to 184, 187 to 193, 198 to 201, 204 to 244, 251 and 252, 255 to 257, 260 to 267, 276 and 277, 280 to 285, 288 to 292, 295 to 297, and 300 to 302, respectively).

Accordingly, in a further aspect of the present invention, there is provided a substantially purified or isolated nucleic acid or nucleic acid fragment including a single nucleotide polymorphism (SNP) from a nucleic acid or nucleic acid fragment according to the present invention or complements or sequences antisense thereto, and functionally active fragments and variants thereof.

In a still further aspect of the present invention there is provided a method of isolating a nucleic acid or nucleic acid fragment of the present invention including a SNP, said method including sequencing nucleic acid fragments from a nucleic acid library.

The nucleic acid library may be of any suitable type and is preferably a cDNA library.

The nucleic acid or nucleic acid fragments may be isolated from a recombinant plasmid or may be amplified, for example using polymerase chain reaction.

The sequencing may be performed by techniques known to those skilled in the art.

In a still further aspect of the present invention, there is provided use of the nucleic acids or nucleic acid fragments of the present invention including SNPs,  
5 and/or nucleotide sequence information thereof, as molecular genetic markers.

In a still further aspect of the present invention there is provided use of a nucleic acid or nucleic acid fragment of the present invention, and/or nucleotide sequence information thereof, as a molecular genetic marker.

More particularly, nucleic acids or nucleic acid fragments according to the  
10 present invention and/or nucleotide sequence information thereof may be used as a molecular genetic marker for quantitative trait loci (QTL) tagging, QTL mapping, DNA fingerprinting and in marker assisted selection, particularly in clovers, alfalfa, ryegrasses and fescues. Even more particularly, nucleic acids or nucleic acid fragments according to the present invention and/or nucleotide sequence  
15 information thereof may be used as molecular genetic markers in plant improvement in relation to plant tolerance to biotic stresses such as viruses, micro-organisms, insects, fungal pathogens; in relation to forage quality; in relation to bloat safety; in relation to condensed tannin content; in relation to plant pigmentation. Even more particularly, sequence information revealing SNPs  
20 allelic variants of the nucleic acids or nucleic acid fragments of the present invention and/or nucleotide sequence information thereof may be used as molecular genetic markers for QTL tagging and mapping and in marker assisted selection, particularly in clovers, alfalfa, ryegrasses and fescues.

In a still further aspect of the present invention there is provided a construct  
25 including a nucleic acid or nucleic acid fragment according to the present invention.

The term "construct" as used herein refers to an artificially assembled or isolated nucleic acid molecule which includes the gene of interest. In general a construct may include the gene or genes of interest, a marker gene which in some

cases can also be the gene of interest and appropriate regulatory sequences. It should be appreciated that the inclusion of regulatory sequences in a construct is optional, for example, such sequences may not be required in situations where the regulatory sequences of a host cell are to be used. The term construct includes  
5 vectors but should not be seen as being limited thereto.

In a still further aspect of the present invention there is provided a vector including a nucleic acid or nucleic acid fragment according to the present invention.

The term "vector" as used herein encompasses both cloning and  
10 expression vectors. Vectors are often recombinant molecules containing nucleic acid molecules from several sources.

In a preferred embodiment of this aspect of the invention, the vector may include a regulatory element such as a promoter, a nucleic acid or nucleic acid fragment according to the present invention and a terminator; said regulatory  
15 element, nucleic acid or nucleic acid fragment and terminator being operatively linked.

By "operatively linked" is meant that said regulatory element is capable of causing expression of said nucleic acid or nucleic acid fragment in a plant cell and said terminator is capable of terminating expression of said nucleic acid or nucleic  
20 acid fragment in a plant cell. Preferably, said regulatory element is upstream of said nucleic acid or nucleic acid fragment and said terminator is downstream of said nucleic acid or nucleic acid fragment.

The vector may be of any suitable type and may be viral or non-viral. The vector may be an expression vector. Such vectors include chromosomal, non-  
25 chromosomal and synthetic nucleic acid sequences, eg. derivatives of plant viruses; bacterial plasmids; derivatives of the Ti plasmid from *Agrobacterium tumefaciens*, derivatives of the Ri plasmid from *Agrobacterium rhizogenes*; phage DNA; yeast artificial chromosomes; bacterial artificial chromosomes; binary bacterial artificial chromosomes; vectors derived from combinations of plasmids

and phage DNA. However, any other vector may be used as long as it is replicable, integrative or viable in the plant cell.

The regulatory element and terminator may be of any suitable type and may be endogenous to the target plant cell or may be exogenous, provided that they  
5 are functional in the target plant cell.

Preferably the regulatory element is a promoter. A variety of promoters which may be employed in the vectors of the present invention are well known to those skilled in the art. Factors influencing the choice of promoter include the desired tissue specificity of the vector, and whether constitutive or inducible  
10 expression is desired and the nature of the plant cell to be transformed (eg. monocotyledon or dicotyledon). Particularly suitable constitutive promoters include the Cauliflower Mosaic Virus 35S (CaMV 35S) promoter and derivatives thereof, the maize Ubiquitin promoter, and the rice Actin promoter.

A variety of terminators which may be employed in the vectors of the  
15 present invention are also well known to those skilled in the art. The terminator may be from the same gene as the promoter sequence or a different gene. Particularly suitable terminators are polyadenylation signals, such as the CaMV 35S polyA and other terminators from the nopaline synthase (*nos*), the octopine synthase (*ocs*) and the *rbcS* genes.

20 The vector, in addition to the regulatory element, the nucleic acid or nucleic acid fragment of the present invention and the terminator, may include further elements necessary for expression of the nucleic acid or nucleic acid fragment, in different combinations, for example vector backbone, origin of replication (*ori*), multiple cloning sites, spacer sequences, enhancers, introns (such as the maize  
25 Ubiquitin *Ubi* intron), antibiotic resistance genes and other selectable marker genes [such as the neomycin phosphotransferase (*npt2*) gene, the hygromycin phosphotransferase (*hph*) gene, the phosphinothricin acetyltransferase (*bar* or *pat*) gene and the gentamycin acetyl transferase (*aacC1*) gene], and reporter genes [such as beta-glucuronidase (*GUS*) gene (*gusA*) and green fluorescent protein



(gfp)]. The vector may also contain a ribosome binding site for translation initiation. The vector may also include appropriate sequences for amplifying expression.

As an alternative to use of a selectable marker gene to provide a phenotypic trait for selection of transformed host cells, the presence of the vector  
5 in transformed cells may be determined by other techniques well known in the art, such as PCR (polymerase chain reaction), Southern blot hybridisation analysis, histochemical GUS assays, northern and Western blot hybridisation analyses.

Those skilled in the art will appreciate that the various components of the vector are operatively linked, so as to result in expression of said nucleic acid or  
10 nucleic acid fragment. Techniques for operatively linking the components of the vector of the present invention are well known to those skilled in the art. Such techniques include the use of linkers, such as synthetic linkers, for example including one or more restriction enzyme sites.

The constructs and vectors of the present invention may be incorporated  
15 into a variety of plants, including monocotyledons (such as grasses from the genera *Lolium*, *Festuca*, *Paspalum*, *Pennisetum*, *Panicum* and other forage and turfgrasses, corn, oat, sugarcane, wheat and barley), dicotyledons (such as *Arabidopsis*, tobacco, clovers, medics, eucalyptus, potato, sugarbeet, canola, soybean, chickpea) and gymnosperms. In a preferred embodiment, the constructs  
20 and vectors may be used to transform monocotyledons, preferably grass species such as ryegrasses (*Lolium* species) and fescues (*Festuca* species), more preferably perennial ryegrass, including forage- and turf-type cultivars. In an alternate preferred embodiment, the constructs and vectors may be used to transform dicotyledons, preferably forage legume species such as clovers  
25 (*Trifolium* species) and medics (*Medicago* species), more preferably white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*) and alfalfa (*Medicago sativa*). Clovers, alfalfa and medics are key pasture legumes in temperate climates throughout the world.

Techniques for incorporating the constructs and vectors of the present  
30 invention into plant cells (for example by transduction, transfection or

transformation) are well known to those skilled in the art. Such techniques include *Agrobacterium* mediated introduction, electroporation to tissues, cells and protoplasts, protoplast fusion, injection into reproductive organs, injection into immature embryos and high velocity projectile introduction to cells, tissues, calli,  
5 immature and mature embryos. The choice of technique will depend largely on the type of plant to be transformed.

Cells incorporating the constructs and vectors of the present invention may be selected, as described above, and then cultured in an appropriate medium to regenerate transformed plants, using techniques well known in the art. The culture  
10 conditions, such as temperature, pH and the like, will be apparent to the person skilled in the art. The resulting plants may be reproduced, either sexually or asexually, using methods well known in the art, to produce successive generations of transformed plants.

In a further aspect of the present invention there is provided a plant cell,  
15 plant, plant seed or other plant part, including, e.g. transformed with, a construct, vector, nucleic acid or nucleic acid fragment of the present invention.

The plant cell, plant, plant seed or other plant part may be from any suitable species, including monocotyledons, dicotyledons and gymnosperms. In a preferred embodiment the plant cell, plant, plant seed or other plant part may be  
20 from a monocotyledon, preferably a grass species, more preferably a ryegrass (*Lolium* species) or fescue (*Festuca* species), even more preferably perennial ryegrass, including both forage- and turf-type cultivars. In an alternate preferred embodiment the plant cell, plant, plant seed or other plant part may be from a dicotyledon, preferably forage legume species such as clovers (*Trifolium* species)  
25 and medics (*Medicago* species), more preferably white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*) and alfalfa (*Medicago sativa*).

The present invention also provides a plant, plant seed or other plant part, or a plant extract derived from a plant cell of the present invention.

The present invention also provides a plant, plant seed or other plant part, or a plant extract derived from a plant of the present invention.

In a further aspect of the present invention there is provided a method of modifying flavonoid biosynthesis in a plant; said method including introducing  
5 into said plant an effective amount of a nucleic acid or nucleic acid fragment, construct and/or a vector according to the present invention.

In a further aspect of the present invention there is provided a method of modifying protein binding, metal chelation, anti-oxidation, and/or UV-light absorption in a plant, said method including introducing into said plant an effective  
10 amount of a nucleic acid or nucleic acid fragment, construct and/or a vector according to the present invention.

In a further aspect of the present invention there is provided a method of modifying pigment production in a plant, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment, construct  
15 and/or a vector according to the present invention.

In a further aspect of the present invention there is provided a method of modifying plant defense to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment, construct and/or a  
20 vector according to the present invention.

In a further aspect of the present invention there is provided a method of modifying forage quality of a plant by disrupting protein foam and/or conferring protection from rumen pasture bloat, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment, construct  
25 and/or a vector according to the present invention.

By "an effective amount" it is meant an amount sufficient to result in an identifiable phenotypic trait in said plant, or a plant, plant seed or other plant part derived therefrom. Such amounts can be readily determined by an appropriately

skilled person, taking into account the type of plant, the route of administration and other relevant factors. Such a person will readily be able to determine a suitable amount and method of administration. See, for example, Maniatis et al, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, the entire disclosure of which is incorporated herein by reference.

Using the methods and materials of the present invention, flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety; and/or isoflavonoid content leading to health benefits, may be increased or otherwise modified, for example by incorporating additional copies of a sense nucleic acid or nucleic acid fragment of the present invention. They may be decreased or otherwise modified, for example by incorporating an antisense nucleic acid or nucleic acid fragment of the present invention.

The present invention will now be more fully described with reference to the accompanying Examples and drawings. It should be understood, however, that the description following is illustrative only and should not be taken in any way as a restriction on the generality of the invention described above.

#### In the Figures

Figure 1 shows the consensus contig nucleotide sequence of TrCH1a (Sequence ID No: 1).

Figure 2 shows the deduced amino acid sequence of TrCH1a (Sequence ID No: 2).

Figure 3 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCH1a (Sequence ID Nos: 3 to 7).

Figure 4 shows the consensus contig nucleotide sequence of TrCH1b (Sequence ID No: 8).

Figure 5 shows the deduced amino acid sequence of TrCH1b (Sequence ID No: 9).

Figure 6 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCH1b (Sequence ID Nos: 10 to 12).

Figure 7 shows the consensus contig nucleotide sequence of TrCH1c (Sequence  
5 ID No: 13).

Figure 8 shows the deduced amino acid sequence of TrCH1c (Sequence ID No: 14).

Figure 9 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCH1c (Sequence ID Nos: 15 and  
10 16).

Figure 10 shows the consensus contig nucleotide sequence of TrCH1d (Sequence ID No: 17).

Figure 11 shows the deduced amino acid sequence of TrCH1d (Sequence ID No: 18).

Figure 12 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCH1d (Sequence ID Nos: 19 to 22).

Figure 13 shows the consensus contig nucleotide sequence of TrCHSa (Sequence ID No: 23).

Figure 14 shows the deduced amino acid sequence of TrCHSa (Sequence ID No: 24).

Figure 15 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSa (Sequence ID Nos: 25 to 63).

Figure 16 shows the consensus contig nucleotide sequence of TrCHSb (Sequence ID No: 64).

Figure 17 shows the deduced amino acid sequence of TrCHSb (Sequence ID No: 65).

- 5 Figure 18 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSb (Sequence ID Nos: 66 to 68).

Figure 19 shows the consensus contig nucleotide sequence of TrCHSc (Sequence ID No: 69).

- 10 Figure 20 shows the deduced amino acid sequence of TrCHSc (Sequence ID No: 70).

Figure 21 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSc (Sequence ID Nos: 71 to 77).

- 15 Figure 22 shows the consensus contig nucleotide sequence of TrCHSd (Sequence ID No: 78).

Figure 23 shows the deduced amino acid sequence of TrCHSd (Sequence ID No: 79).

- 20 Figure 24 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSd (Sequence ID Nos: 80 to 90).

Figure 25 shows the consensus contig nucleotide sequence of TrCHSe (Sequence ID No: 91).

- 25 Figure 26 shows the deduced amino acid sequence of TrCHSe (Sequence ID No: 92).

Figure 27 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSe (Sequence ID Nos: 93 and 94).

Figure 28 shows the consensus contig nucleotide sequence of TrCHSf (Sequence  
5 ID No: 95).

Figure 29 shows the deduced amino acid sequence of TrCHSf (Sequence ID No: 96).

Figure 30 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSf (Sequence ID Nos: 97 to  
10 100).

Figure 31 shows the consensus contig nucleotide sequence of TrCHSg (Sequence ID No: 101).

Figure 32 shows the deduced amino acid sequence of TrCHSg (Sequence ID No: 102).

15 Figure 33 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHSg (Sequence ID Nos: 103 to 105).

Figure 34 shows the consensus contig nucleotide sequence of TrCHSh (Sequence ID No: 106).

20 Figure 35 shows the deduced amino acid sequence of TrCHSh (Sequence ID No: 107).

Figure 36 shows the nucleotide sequence of TrCHRa (Sequence ID No: 108).

Figure 37 shows the deduced amino acid sequence of TrCHRa (Sequence ID No: 109).

Figure 38 shows the consensus contig nucleotide sequence of TrCHRB (Sequence ID No: 110).

Figure 39 shows the deduced amino acid sequence of TrCHRB (Sequence ID No: 111).

- 5 Figure 40 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHRB (Sequence ID Nos: 112 to 116).

Figure 41 shows the consensus contig nucleotide sequence of TrCHRC (Sequence ID No: 117).

- 10 Figure 42 shows the deduced amino acid sequence of TrCHRC (Sequence ID No: 118).

Figure 43 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrCHRC (Sequence ID Nos: 119 to 134).

- 15 Figure 44 shows the consensus contig nucleotide sequence of TrDFRa (Sequence ID No: 135).

Figure 45 shows the deduced amino acid sequence of TrDFRa (Sequence ID No: 136).

- 20 Figure 46 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrDFRa (Sequence ID Nos: 137 to 146).

Figure 47 shows the consensus contig nucleotide sequence of TrDFRb (Sequence ID No: 147).

- 25 Figure 48 shows the deduced amino acid sequence of TrDFRb (Sequence ID No: 148).



Figure 49 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrDFRb (Sequence ID Nos: 149 to 152).

Figure 50 shows the nucleotide sequence of TrDFRc (Sequence ID No: 153).

- 5 Figure 51 shows the deduced amino acid sequence of TrDFRc (Sequence ID No: 154).

Figure 52 shows the consensus contig nucleotide sequence of TrDFRd (Sequence ID No: 155).

- 10 Figure 53 shows the deduced amino acid sequence of TrDFRd (Sequence ID No: 156).

Figure 54 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrDFRd (Sequence ID Nos: 157 and 158).

Figure 55 shows the nucleotide sequence of TrDFRe (Sequence ID No: 159).

- 15 Figure 56 shows the deduced amino acid sequence of TrDFRe (Sequence ID No: 160).

Figure 57 shows the nucleotide sequence of TrDFRf (Sequence ID No: 161).

Figure 58 shows the deduced amino acid sequence of TrDFRf (Sequence ID No: 162).

- 20 Figure 59 shows the consensus contig nucleotide sequence of TrDFRg (Sequence ID No: 163).

Figure 60 shows the deduced amino acid sequence of TrDFRg (Sequence ID No: 164).

Figure 61 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrDFRg (Sequence ID Nos: 165 to 167).

Figure 62 shows the consensus contig nucleotide sequence of TrDFRh (Sequence  
5 ID No: 168).

Figure 63 shows the deduced amino acid sequence of TrDFRh (Sequence ID No: 169).

Figure 64 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrDFRh (Sequence ID Nos: 170 to  
10 184).

Figure 65 shows the consensus contig nucleotide sequence of TrLCRa (Sequence ID No: 185).

Figure 66 shows the deduced amino acid sequence of TrLCRa (Sequence ID No: 186).

Figure 67 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrLCRa (Sequence ID Nos: 187 to  
15 193).

Figure 68 shows the nucleotide sequence of TrF3'5'Ha (Sequence ID No: 194).

Figure 69 shows the deduced amino acid sequence of TrF3'5'Ha (Sequence ID  
20 No: 195).

Figure 70 shows the consensus contig nucleotide sequence of TrF3'5'Hb (Sequence ID No: 196).

Figure 71 shows the deduced amino acid sequence of TrF3'5'Hb (Sequence ID No: 197).

Figure 72 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrF3'5'Hb (Sequence ID Nos: 198 to 201).

Figure 73 shows the consensus contig nucleotide sequence of TrF3Ha (Sequence  
5 ID No: 202).

Figure 74 shows the deduced amino acid sequence of TrF3Ha (Sequence ID No: 203).

Figure 75 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrF3Ha (Sequence ID Nos: 204 to  
10 244).

Figure 76 shows the nucleotide sequence of TrF3Hb (Sequence ID No: 245).

Figure 77 shows the deduced amino acid sequence of TrF3Hb (Sequence ID No: 246).

Figure 78 shows the nucleotide sequence of TrF3Hc (Sequence ID No: 247).

15 Figure 79 shows the deduced amino acid sequence of TrF3Hc (Sequence ID No: 248).

Figure 80 shows the consensus contig nucleotide sequence of TrF3'Ha (Sequence ID No: 249).

Figure 81 shows the deduced amino acid sequence of TrF3'Ha (Sequence ID No:  
20 250).

Figure 82 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrF3'Ha (Sequence ID Nos: 251 and 252).

Figure 83 shows the consensus contig nucleotide sequence of TrPALa (Sequence ID No: 253).

Figure 84 shows the deduced amino acid sequence of TrPALa (Sequence ID No: 254).

- 5 Figure 85 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrPALa (Sequence ID Nos: 255 to 257).

Figure 86 shows the consensus contig nucleotide sequence of TrPALb (Sequence ID No: 258).

- 10 Figure 87 shows the deduced amino acid sequence of TrPALb (Sequence ID No: 259).

Figure 88 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrPALb (Sequence ID Nos: 260 to 267).

- 15 Figure 89 shows the nucleotide sequence of TrPALc (Sequence ID No: 268).

Figure 90 shows the deduced amino acid sequence of TrPALc (Sequence ID No: 269).

Figure 91 shows the nucleotide sequence of TrPALd (Sequence ID No: 270).

- Figure 92 shows the deduced amino acid sequence of TrPALd (Sequence ID No:  
20 271).

Figure 93 shows the nucleotide sequence of TrPALe (Sequence ID No: 272).

Figure 94 shows the deduced amino acid sequence of TrPALe (Sequence ID No: 273).

Figure 95 shows the consensus contig nucleotide sequence of TrPALf (Sequence ID No: 274).

Figure 96 shows the deduced amino acid sequence of TrPALf (Sequence ID No: 275).

- 5 Figure 97 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrPALf (Sequence ID Nos: 276 and 277).

Figure 98 shows the consensus contig nucleotide sequence of TrVRa (Sequence ID No: 278).

- 10 Figure 99 shows the deduced amino acid sequence of TrVRa (Sequence ID No: 279).

Figure 100 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence TrVRa (Sequence ID Nos: 280 to 285).

- 15 Figure 101 shows the consensus contig nucleotide sequence of LpDFRa (Sequence ID No: 286).

Figure 102 shows the deduced amino acid sequence of LpDFRa (Sequence ID No: 287).

- 20 Figure 103 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence LpDFRa (Sequence ID Nos: 288 to 292).

Figure 104 shows the consensus contig nucleotide sequence of LpDFRb (Sequence ID No: 293).

- 25 Figure 105 shows the deduced amino acid sequence of LpDFRb (Sequence ID No: 294).

Figure 106 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence LpDFRb (Sequence ID Nos: 295 to 297).

Figure 107 shows the consensus contig nucleotide sequence of LpF3Ha  
5 (Sequence ID No: 298).

Figure 108 shows the deduced amino acid sequence of LpF3Ha (Sequence ID No: 299).

Figure 109 shows the nucleotide sequences of the nucleic acid fragments contributing to the consensus contig sequence LpF3Ha (Sequence ID Nos: 300 to  
10 302).

Figure 110 shows a plasmid map of the cDNA encoding perennial ryegrass F3OH.

Figure 111 shows the full nucleotide sequence of perennial ryegrass F3OH cDNA (Sequence ID No: 303).

Figure 112 shows the deduced amino acid sequence of perennial ryegrass F3OH  
15 cDNA (Sequence ID No: 304).

Figure 113 shows plasmid maps of sense and antisense constructs of LpF3OH in pDH51 transformation vector.

Figure 114 shows plasmid maps of sense and antisense constructs of LpF3OH in pPZP221:35S<sup>2</sup> binary transformation vector.

20 Figure 115 shows screening by Southern hybridisation for RFLPs using LpF3OH as a probe.

Figure 116 shows a plasmid map of the cDNA encoding white clover BANa.

Figure 117 shows the full nucleotide sequence of white clover BANa cDNA (Sequence ID No: 305).

Figure 118 shows the deduced amino acid sequence of white clover BANa cDNA (Sequence ID No: 306).

Figure 119 shows plasmid maps of sense and antisense constructs of TrBANa in pDH51 transformation vector.

- 5 Figure 120 shows plasmid maps of sense and antisense constructs of TrBANa in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 121 shows a plasmid map of the cDNA encoding white clover CH1a.

Figure 122 shows the full nucleotide sequence of white clover CH1a cDNA (Sequence ID No: 307).

- 10 Figure 123 shows the deduced amino acid sequence of white clover CH1a cDNA (Sequence ID No: 308).

Figure 124 shows plasmid maps of sense and antisense constructs of TrCH1a in pDH51 transformation vector.

- 15 Figure 125 shows plasmid maps of sense and antisense constructs of TrCH1a in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 126 shows a plasmid map of the cDNA encoding white clover CH1d.

Figure 127 shows the full nucleotide sequence of white clover CH1d cDNA (Sequence ID No: 309).

- 20 Figure 128 shows the deduced amino acid sequence of white clover CH1d cDNA (Sequence ID No: 310).

Figure 129 shows plasmid maps of sense and antisense constructs of TrCH1d in pDH51 transformation vector.

Figure 130 shows plasmid maps of sense and antisense constructs of TrCHId in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 131 shows a plasmid map of the cDNA encoding white clover CHRc.

Figure 132 shows the full nucleotide sequence of white clover CHRc cDNA  
5 (Sequence ID No: 311).

Figure 133 shows the deduced amino acid sequence of white clover CHRc cDNA (Sequence ID No: 312).

Figure 134 shows plasmid maps of sense and antisense constructs of TrCHRc in pDH51 transformation vector.

10 Figure 135 shows plasmid maps of sense and antisense constructs of TrCHRc in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 136 shows a plasmid map of the cDNA encoding white clover CHSa1.

Figure 137 shows the full nucleotide sequence of white clover CHSa1 cDNA (Sequence ID No: 313).

15 Figure 138 shows the deduced amino acid sequence of white clover CHSa1 cDNA (Sequence ID No: 314).

Figure 139 shows plasmid maps of sense and antisense constructs of TrCHSa1 in pDH51 transformation vector.

Figure 140 shows plasmid maps of sense and antisense constructs of TrCHSa1  
20 pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 141 shows a plasmid map of the cDNA encoding white clover CHSa3.

Figure 142 shows the full nucleotide sequence of white clover CHSa3 cDNA (Sequence ID No: 315).



Figure 143 shows the deduced amino acid sequence of white clover CHSa3 cDNA (Sequence ID No: 316).

Figure 144 shows plasmid maps of sense and antisense constructs of TrCHSa3 in pDH51 transformation vector.

- 5 Figure 145 shows plasmid maps of sense and antisense constructs of TrCHSa3 in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 146 shows a plasmid map of the cDNA encoding white clover CHSc.

Figure 147 shows the full nucleotide sequence of white clover CHSc cDNA (Sequence ID No: 317).

- 10 Figure 148 shows the deduced amino acid sequence of white clover CHSc cDNA (Sequence ID No: 318).

Figure 149 shows plasmid maps of sense and antisense constructs of TrCHSc in pDH51 transformation vector.

- Figure 150 shows plasmid maps of sense and antisense constructs of TrCHSc in  
15 pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 151 shows a plasmid map of the cDNA encoding white clover CHSd2.

Figure 152 shows the full nucleotide sequence of white clover CHSd2 cDNA (Sequence ID No: 319).

- Figure 153 shows the deduced amino acid sequence of white clover CHSd2 cDNA  
20 (Sequence ID No: 320).

Figure 154 shows plasmid maps of sense and antisense constructs of TrCHSd2 in pDH51 transformation vector.

Figure 155 shows plasmid maps of sense and antisense constructs of TrCHSd2 in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 156 shows a plasmid map of the cDNA encoding white clover CHSf.

Figure 157 shows the full nucleotide sequence of white clover CHSf cDNA  
5 (Sequence ID No: 321).

Figure 158 shows the deduced amino acid sequence of white clover CHSf cDNA (Sequence ID No: 322).

Figure 159 shows plasmid maps of sense and antisense constructs of TrCHSf in pDH51 transformation vector.

10 Figure 160 shows plasmid maps of sense and antisense constructs of TrCHSf in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 161 shows a plasmid map of the cDNA encoding white clover CHSh.

Figure 162 shows the full nucleotide sequence of white clover CHSh cDNA (Sequence ID No: 323).

15 Figure 163 shows the deduced amino acid sequence of white clover CHSh cDNA (Sequence ID No: 324).

Figure 164 shows plasmid maps of sense and antisense constructs of TrCHSh in pDH51 transformation vector.

Figure 165 shows plasmid maps of sense and antisense constructs of TrCHSh in  
20 pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 166 shows a plasmid map of the cDNA encoding white clover DFRd.

Figure 167 shows the full nucleotide sequence of white clover DFRd cDNA (Sequence ID No: 325).

Figure 168 shows the deduced amino acid sequence of white clover DFRd cDNA (Sequence ID No: 326).

Figure 169 shows plasmid maps of sense and antisense constructs of TrDFRd in pDH51 transformation vector.

- 5 Figure 170 shows plasmid maps of sense and antisense constructs of TrDFRd in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 171 shows a plasmid map of the cDNA encoding white clover F3Ha.

Figure 172 shows the full nucleotide sequence of white clover F3Ha cDNA (Sequence ID No: 327).

- 10 Figure 173 shows the deduced amino acid sequence of white clover F3Ha cDNA (Sequence ID No: 328).

Figure 174 shows plasmid maps of sense and antisense constructs of TrF3Ha in pDH51 transformation vector.

- 15 Figure 175 shows plasmid maps of sense and antisense constructs of TrF3Ha in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 176 shows a plasmid map of the cDNA encoding white clover PALa.

Figure 177 shows the full nucleotide sequence of white clover PALa cDNA (Sequence ID No: 329).

- 20 Figure 178 shows the deduced amino acid sequence of white clover PALa cDNA (Sequence ID No: 330).

Figure 179 shows plasmid maps of sense and antisense constructs of TrPALa in pDH51 transformation vector.

Figure 180 shows plasmid maps of sense and antisense constructs of TrPALa in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 181 shows a plasmid map of the cDNA encoding white clover PALb.

Figure 182 shows the full nucleotide sequence of white clover PALb cDNA  
5 (Sequence ID No: 331).

Figure 183 shows the deduced amino acid sequence of white clover PALb cDNA (Sequence ID No: 332).

Figure 184 shows plasmid maps of sense and antisense constructs of TrPALb in pDH51 transformation vector.

10 Figure 185 shows plasmid maps of sense and antisense constructs of TrPALb in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 186 shows a plasmid map of the cDNA encoding white clover PALf.

Figure 187 shows the full nucleotide sequence of white clover PALf cDNA (Sequence ID No: 333).

15 Figure 188 shows the deduced amino acid sequence of white clover PALf cDNA (Sequence ID No: 334).

Figure 189 shows plasmid maps of sense and antisense constructs of TrPALf in pDH51 transformation vector.

Figure 190 shows plasmid maps of sense and antisense constructs of TrPALf in  
20 pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 191 shows a plasmid map of the cDNA encoding white clover VRa.

Figure 192 shows the full nucleotide sequence of white clover VRa cDNA (Sequence ID No: 335).

Figure 193 shows the deduced amino acid sequence of white clover VRa cDNA (Sequence ID No: 336).

Figure 194 shows plasmid maps of sense and antisense constructs of TrVRa in pDH51 transformation vector.

- 5 Figure 195 shows plasmid maps of sense and antisense constructs of TrVRa in pPZP221:35S<sup>2</sup> binary transformation vector.

Figure 196 shows A, infiltration of Arabidopsis plants; B, selection of transgenic Arabidopsis plants on medium containing 75 µg/ml gentamycin; C, young transgenic Arabidopsis plants; D, E, two representative results of real-time PCR  
10 analysis of Arabidopsis transformed with chimeric genes involved in flavonoid biosynthesis.

Figure 197 shows the genetic map detailing the relation of perennial ryegrass genes involved in flavonoid biosynthesis.

### EXAMPLE 1

- 15 **Preparation of cDNA libraries, isolation and sequencing of cDNAs coding for CHI, CHI-like, CHS, CHS-like, CHR, CHR-like, DFR, DFR-like, LCR, LCR-like, F3'5'H, F3'5'H-like, F3H, F3H-like, F3'H, F3'H-like, PAL, PAL-like, VR and VR-like proteins from white clover (*Trifolium repens*) and perennial ryegrass (*Lolium perenne*)**

20

cDNA libraries representing mRNAs from various organs and tissues of white clover (*Trifolium repens*) and perennial ryegrass (*Lolium perenne*) were prepared. The characteristics of the white clover and perennial ryegrass libraries, respectively, are described below (Tables 1 and 2).

**TABLE 1****cDNA libraries from white clover (*Trifolium repens*)**

<b>Library</b>	<b>Organ/Tissue</b>
01wc	Whole seedling, light grown
02wc	Nodulated root 3, 5, 10, 14, 21 & 28 day old seedling
03wc	Nodules pinched off roots of 42 day old rhizobium inoculated plants
04wc	Cut leaf and stem collected after 0, 1, 4, 6 & 14 h after cutting
05wc	Inflorescences: <50% open, not fully open and fully open
06wc	Dark grown etiolated
07wc	Inflorescence – very early stages, stem elongation, < 15 petals, 15-20 petals
08wc	seed frozen at –80°C, imbibed in dark overnight at 10°C
09wc	Drought stressed plants
10wc	AMV infected leaf
11wc	WCMV infected leaf
12wc	Phosphorus starved plants
13wc	Vegetative stolon tip
14wc	stolon root initials
15wc	Senescing stolon
16wc	Senescing leaf

**TABLE 2**

5

**cDNA libraries from perennial ryegrass (*Lolium perenne*)**

<b>Library</b>	<b>Organ/Tissue</b>
01rg	Roots from 3-4 day old light-grown seedlings
02rg	Leaves from 3-4 day old light-grown seedlings
03rg	Etiolated 3-4 day old dark-grown seedlings
04rg	Whole etiolated seedlings (1-5 day old and 17 days old)
05rg	Senescing leaves from mature plants

Library	Organ/Tissue
06rg	Whole etiolated seedlings (1-5 day old and 17 days old)
07rg	Roots from mature plants grown in hydroponic culture
08rg	Senescent leaf tissue
09rg	Whole tillers and sliced leaves (0, 1, 3, 6, 12 and 24 h after harvesting)
10rg	Embryogenic suspension-cultured cells
11rg	Non-embryogenic suspension-cultured cells
12rg	Whole tillers and sliced leaves (0, 1, 3, 6, 12 and 24 h after harvesting)
13rg	Shoot apices including vegetative apical meristems
14rg	Immature inflorescences including different stages of inflorescence meristem and inflorescence development
15rg	Defatted pollen
16rg	Leaf blades and leaf sheaths ( <i>rbcL</i> , <i>rbcS</i> , <i>cab</i> , <i>wir2A</i> subtracted)
17rg	Senescing leaves and tillers
18rg	Drought-stressed tillers (pseudostems from plants subjected to PEG-simulated drought stress)
19rg	Non-embryogenic suspension-cultured cells subjected to osmotic stress (grown in media with half-strength salts) (1, 2, 3, 4, 5, 6, 24 and 48 h after transfer)
20rg	Non-embryogenic suspension-cultured cells subjected to osmotic stress (grown in media with double-strength salts) (1, 2, 3, 4, 5, 6, 24 and 48 h after transfer)
21rg	Drought-stressed tillers (pseudostems from plants subjected to PEG-simulated drought stress)
22rg	Spikelets with open and maturing florets
23rg	Mature roots (specific subtraction with leaf tissue)

The cDNA libraries may be prepared by any of many methods available. For example, total RNA may be isolated using the Trizol method (Gibco-BRL, USA) or the RNeasy Plant Mini kit (Qiagen, Germany), following the

5 manufacturers' instructions. cDNAs may be generated using the SMART PCR

- cDNA synthesis kit (Clontech, USA), cDNAs may be amplified by long distance polymerase chain reaction using the Advantage 2 PCR Enzyme system (Clontech, USA), cDNAs may be cleaned using the GeneClean spin column (Bio 101, USA), tailed and size fractionated, according to the protocol provided by Clontech. The
- 5 cDNAs may be introduced into the pGEM-T Easy Vector system 1 (Promega, USA) according to the protocol provided by Promega. The cDNAs in the pGEM-T Easy plasmid vector are transfected into *Escherichia coli* Epicurian coli XL10-Gold ultra competent cells (Stratagene, USA) according to the protocol provided by Stratagene.
- 10 Alternatively, the cDNAs may be introduced into plasmid vectors for first preparing the cDNA libraries in Uni-ZAP XR vectors according to the manufacturer's protocol (Stratagene Cloning Systems, La Jolla, CA, USA). The Uni-ZAP XR libraries are converted into plasmid libraries according to the protocol provided by Stratagene. Upon conversion, cDNA inserts will be contained in the
- 15 plasmid vector pBluescript. In addition, the cDNAs may be introduced directly into precut pBluescript II SK(+) vectors (Stratagene) using T4 DNA ligase (New England Biolabs), followed by transfection into *E. coli* DH10B cells according to the manufacturer's protocol (GIBCO BRL Products).
- Once the cDNA inserts are in plasmid vectors, plasmid DNAs are prepared
- 20 from randomly picked bacterial colonies containing recombinant plasmids, or the insert cDNA sequences are amplified via polymerase chain reaction using primers specific for vector sequences flanking the inserted cDNA sequences. Plasmid DNA preparation may be performed robotically using the Qiagen QiaPrep Turbo kit (Qiagen, Germany) according to the protocol provided by Qiagen. Amplified insert
- 25 DNAs are sequenced in dye-terminator sequencing reactions to generate partial cDNA sequences (expressed sequence tags or "ESTs"). The resulting ESTs are analyzed using an Applied Biosystems ABI 3700 sequence analyser.



## EXAMPLE 2

### DNA sequence analyses

The cDNA clones encoding CHI, CHI-like, CHS, CHS-like, CHR, CHR-like, DFR, DFR-like, LCR, LCR-like, F3'5'H, F3'5'H-like, F3H, F3H-like, F3'H, F3'H-like, PAL, PAL-like, VR and VR-like proteins were identified by conducting BLAST (Basic Local Alignment Search Tool; Altschul *et al.* (1993) *J. Mol. Biol.* 215:403-410) searches. The cDNA sequences obtained were analysed for similarity to all publicly available DNA sequences contained in the eBioinformatics nucleotide database using the BLASTN algorithm provided by the National Center for Biotechnology Information (NCBI). The DNA sequences were translated in all reading frames and compared for similarity to all publicly available protein sequences contained in the SWISS-PROT protein sequence database using BLASTx algorithm (v 2.0.1) (Gish and States (1993) *Nature Genetics* 3:266-272) provided by the NCBI.

The cDNA sequences obtained and identified were then used to identify additional identical and/or overlapping cDNA sequences generated using the BLASTN algorithm. The identical and/or overlapping sequences were subjected to a multiple alignment using the CLUSTALw algorithm, and to generate a consensus contig sequence derived from this multiple sequence alignment. The consensus contig sequence was then used as a query for a search against the SWISS-PROT protein sequence database using the BLASTx algorithm to confirm the initial identification.

## EXAMPLE 3

**Identification and full-length sequencing of cDNAs encoding perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa proteins**

To fully characterise for the purposes of the generation of probes for hybridisation experiments and the generation of transformation vectors, a set of

cDNAs encoding perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa proteins was identified and fully sequenced.

Full-length cDNAs were identified from our EST sequence database using  
5 relevant published sequences (NCBI databank) as queries for BLAST searches. Full-length cDNAs were identified by alignment of the query and hit sequences using Sequencher (Gene Codes Corp., Ann Arbor, MI 48108, USA). The original plasmid was then used to transform chemically competent XL-1 cells (prepared in-house, CaCl<sub>2</sub> protocol). After colony PCR (using HotStarTaq, Qiagen) a minimum  
10 of three PCR-positive colonies per transformation were picked for initial sequencing with M13F and M13R primers. The resulting sequences were aligned with the original EST sequence using Sequencher to confirm identity and one of the three clones was picked for full-length sequencing, usually the one with the best initial sequencing result.

15 Sequencing was completed by primer walking, i.e. oligonucleotide primers were designed to the initial sequence and used for further sequencing. In most cases the sequencing could be done from both 5' and 3' end. The sequences of the oligonucleotide primers are shown in Table 2. In some instances, however, an extended poly-A tail necessitated the sequencing of the cDNA to be completed  
20 from the 5' end.

Contigs were then assembled in Sequencher. The contigs include the sequences of the SMART primers used to generate the initial cDNA library as well as pGEM-T Easy vector sequence up to the EcoRI cut site both at the 5' and 3' end.

25 Plasmid maps and the full cDNA sequences of perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa proteins were obtained (Figures 110, 116, 121, 126, 131, 136, 141, 146, 151, 156, 161, 166, 171, 176, 181, 186 and 191).

TABLE 2

List of primers used for sequencing of the full-length cDNAs

gene name	clone ID	sequencing primer	primer sequence (5'>3')
LpF3OH	08rg1YsF07	08rg1YsF07.f1	TTGAGAGCTTCGTCGACC
		08rg1YsF07.r1	AACTCCTCGTAGTACTCC
TrCHRe	11wc1IsD03	11wc1IsD03.f1	TTCAATTGGAGTACTTGG
		11wc1IsD03.r1	ACTCCTTGTTTCATATAACC
TrCHSa1	02wc2FsD07	02wc2FsD07.f1	ACATGGTGGTGGTTGAGG
		02wc2FsD07.f2	TGCTGCACTCATTGTTGG
		02wc2FsD07.f3	ACATTGATAAGGCATTGG
TrCHSa3	05wc1RsB06	05wc1RsB06.f1	AGGAGGCTGCAGTCAAGG
		05wc1RsB06.f2	TGCCTGAAATTGAGAAACC
		05wc1RsB06.f3	AAAGCTAGCCTTGAAGCC
TrCHSc	07wc1TsE12	07wc1TsE12.f1	TCGGACATAACTCATGTGG
		07wc1TsE12.f2	TTGGGTTGGAGAATAAGG
		07wc1TsE12.r1	TGGACATTTATTGGTTGC
		07wc1TsE12.r2	TATCATGTCTGGAAATGC
TrCHSd2	07wc1XsD03	07wc1XsD03.f1	TTTATGTGAGTACATGGC
		07wc1XsD03.f2	AGCAGCTGTGATTGTAGG
		07wc1XsD03.f3	TGAGAAAGCTCTTGTGAGG
TrCHSf	07wc1UsD07	07wc1UsD07.f1	AGATTGCATCAAAGAATGG
		07wc1UsD07.r1	GGTCCAAAAGCCAATCC
TrCHSh	13wc2IsG04	13wc2IsG04.f1	TAAGACGAGACATAGTGG
		13wc2IsG04.r1	TATTCAC TAAGCACATGC
TrDFRd	12wc1CsE09	12wc1CsE09.f1	TTACCTCGTCTGTCTCG
		12wc1CsE09.r1	AACACACACATGTCTACC
TrF3Ha	07wc1LsG03	07wc1LsG03.f1	TGAAGGATTGGAGAGAGC
		07wc1LsG03.r1	TACACAGTTGCATCTGG
TrPALa	04wc1UsB03	04wc1UsB03.f1	ATCGGAATCTGCTAGAGC
		04wc1UsB03.f2	TGTTGGTTCTGGTTTAGC
		04wc1UsB03.r1	TTCATATGCAATCCTTGC
		04wc1UsB03.r2	TCTTGGTTGTGTTGTTCC

TrPALb	05wc1PsH02	05wc1PsH02.f1	TGGGACTGATAGTTATGG
		05wc1PsH02.f2	TCTTGCTCTTGTTAATGG
		05wc1PsH02.r1	AGCACCATTCCACTCTCC
		05wc1PsH02.r2	TTCTCTTCGCTACTTGGC
TrPALf	13wc2AsD12	13wc2AsD12.f1	ATAGTGGTGTGAGGGTGG
		13wc2AsD12.f2	TCTTGTTAATGGTACTGC
		13wc2AsD12.r1	ATTTATCGCACTCTTCGC
		13wc2AsD12.r2	AAAGTGGGAAGACATGAGC
TrVRa	11wc1NsA07	11wc1NsA07.f1	AAGAACAGTGGATGGAGC
		11wc1NsA07.r1	TCAACTCATCTACTGATAG

#### EXAMPLE 4

**Development of transformation vectors containing chimeric genes with cDNA sequences from perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa**

To alter the expression of the proteins involved in flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits, perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa, through antisense and/or sense suppression technology and for over-expression of these key enzymes in transgenic plants, a set of sense and antisense transformation vectors was produced.

cDNA fragments were generated by high fidelity PCR using the original pGEM-T Easy plasmid cDNA as a template. The primers used (Table 3) contained recognition sites for appropriate restriction enzymes, for example EcoRI and XbaI, for directional and non-directional cloning into the target vector. After PCR amplification and restriction digest with the appropriate restriction enzyme (usually

XbaI), the cDNA fragments were cloned into the corresponding site in pDH51, a pUC18-based transformation vector containing a CaMV 35S expression cassette. The orientation of the constructs (sense or antisense) was checked by DNA sequencing through the multi-cloning site of the vector. Transformation vectors

5 containing chimeric genes using full-length open reading frame cDNAs encoding perennial ryegrass F3OH and white clover BANA, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa proteins in sense and antisense orientations under the control of the CaMV 35S promoter were generated (Figures 113, 119, 124, 129, 134, 139, 144, 149, 154,

10 159, 164, 169, 174, 179, 184, 189 and 194).

**TABLE 3****List of primers used to PCR-amplify the open reading frames**

gene name	clone ID	primer	primer sequence (5'→3')
LpF3OH	08rg1YsF07	08rg1YsF07f	GAATTCTAGAAGCAGAAAGTACGGACATCAGC
		08rg1YsF07r	GAATTCTAGAACCATATGGCGACACATCG
TrBANA	05wc2XsG02	05wc2XsG02f	GGATCCTCTAGAGCACTAGTGTGTATAAGTTTCTT GG
		05wc2XsG02r	GGATCCTCTAGACCCCTTAGTCTTAAATACTCG
TrCH1a	06wc2AsF12	06wc2AsF12f	GAATTCTAGAGATCTGAAACAACATAGTCACC
		06wc2AsF12r	GAATTCTAGATCAATCTTGTGCTGCAATGC
TrCH1d	12wc1FsG04	12wc1FsG04f	GAATTCTAGAAAGTTCAACGAGATCAATGG
		12wc1FsG04r	GAATTCTAGATTCCGCTTGGTCTTTATTGC
TrCHRc	11wc1IsD03	11wc1IsD03f	GAATTCTAGAACATGGGTAGTGTGAAATTCC
		11wc1IsD03r	GAATTCTAGAAGATATTGAGTGAGCTTAAGG
TrCHSa1	02wc2FsD07	02wc2FsD07f	GACGTCGACATTACATACATAGCAGGAAC
		02wc2FsD07r	GACGTCGACAGTCTCTCATTCTCATATAGC
TrCHSa3	05wc1RsB06	05wc1RsB06f	GAATTCTAGAAGATATGGTGAGGTAGCTG
		05wc1RsB06r	GAATTCTAGAATCACACATCTTATATAGCC
TrCHSc	07wc1TsE12	07wc1TsE12f	GAATTCTAGAAGAAGAAATATGGGAGACGAAGG
		07wc1TsE12r	GAATTCTAGAAAGACTTCATGCACACAAGTTCC
TrCHSd2	07wc1XsD03	07wc1XsD03f	GAATTCTAGAATAACCTATCAGTACTCACC
		07wc1XsD03r	GAATTCTAGAATCTAGGCAATTTAAGTGCC

TrCHSf	07wc1UsD07	07wc1UsD07f	GAATTCTAGATGATTCATTGTTGTTTCCATAAC
		07wc1UsD07r	GAATTCTAGAACATATTCATCTTCCTATCAC
TrCHSh	13wc2IsG04	13wc2IsG04f	GAATTCTAGATCCAAATTCTCGTACCTCACC
		13wc2IsG04r	GAATTCTAGATAGTTCACATCTCTCGGCAGG
TrDFRd	12wc1CsE09	12wc1CsE09f	GACGTCGACACAACAGTCTTCCACTTGAGC
		12wc1CsE09r	GACGTCGACTCTATACTCTGGTAACTATAGG
TrF3Ha	07wc1LsG03	07wc1LsG03f	GAATTCTAGAACCACACAACACACAAACACC
		07wc1LsG03r	GAATTCTAGAACCAAGCAGCTTAATACACG
TrPALa	04wc1UsB03	04wc1UsB03f	AGTACTGCAGAGATATGGAAGTAGTAGCAGCAGC
		04wc1UsB03r	AGTACTGCAGTAGCAAACCAGTTCCCAACTCC
TrPALb	05wc1PsH02	05wc1PsH02f	AGTACTGCAGATAATGGAGGGAATTACCAATGG
		05wc1PsH02r	AGTACTGCAGTGCTAATTAACATATTGGTAGAGG
TrPALf	13wc2AsD12	13wc2AsD12f	AGTACTGCAGATAATGGAGGGAATTACCAATGG
		13wc2AsD12r	AGTACTGCAGTGCTAATTAACATATTGGTAGAGG
TrVRa	11wc1NsA07	11wc1NsA07f	AGTACTGCAGATAAAGAGAGTCAAAAATGGC
		11wc1NsA07r	AGTACTGCAGAACACATACTTAGAGATAGCC

### EXAMPLE 5

**Development of binary transformation vectors containing chimeric genes with cDNA sequences from perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa**

To alter the expression of the proteins involved in flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits, perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa, through antisense and/or sense suppression technology and for over-expression of these key proteins in transgenic plants, a set of sense and antisense binary transformation vectors was produced.

cDNA fragments were generated by high fidelity PCR using the original pGEM-T Easy plasmid cDNA as a template. The primers used (Table 3) contained recognition sites for appropriate restriction enzymes, for example EcoRI and XbaI, for directional and non-directional cloning into the target vector. After PCR  
5 amplification and restriction digest with the appropriate restriction enzyme (usually XbaI), the cDNA fragments were cloned into the corresponding site in a modified pPZP binary vector (Hajdukiewicz *et al.*, 1994). The pPZP221 vector was modified to contain the 35S<sup>2</sup> cassette from pKYLX71:35S<sup>2</sup> as follows. pKYLX71:35S<sup>2</sup> was cut with ClaI. The 5' overhang was filled in using Klenow and the blunt end was A-  
10 tailed with Taq polymerase. After cutting with EcoRI, the 2kb fragment with an EcoRI-compatible and a 3'-A tail was gel-purified. pPZP221 was cut with HindIII and the resulting 5' overhang filled in and T-tailed with Taq polymerase. The remainder of the original pPZP221 multi-cloning site was removed by digestion with EcoRI, and the expression cassette cloned into the EcoRI site and the 3' T  
15 overhang restoring the HindIII site. This binary vector contains between the left and right border the plant selectable marker gene *aaaC1* under the control of the 35S promoter and 35S terminator and the pKYLX71:35S<sup>2</sup>-derived expression cassette with a CaMV 35S promoter with a duplicated enhancer region and an *rbcS* terminator.

20 The orientation of the constructs (sense or antisense) was checked by restriction enzyme digest. Transformation vectors containing chimeric genes using full-length open reading frame cDNAs encoding perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa proteins in sense and antisense  
25 orientations under the control of the CaMV 35S<sup>2</sup> promoter were generated (Figures 114, 120, 125, 130, 135, 140, 145, 150, 155, 160, 165, 170, 175, 180, 185, 190 and 195).

**EXAMPLE 6**

**Production and analysis of transgenic Arabidopsis plants carrying chimeric perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa genes involved in flavonoid biosynthesis**

A set of transgenic Arabidopsis plants carrying chimeric perennial ryegrass and white clover genes involved in flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits, were produced.

pPZP221-based transformation vectors with *LpF3OH* and *TrBANa*, *TrCH1a*, *TrCH1d*, *TrCHRc*, *TrCHSa1*, *TrCHSa3*, *TrCHSc*, *TrCHSd2*, *TrCHSf*, *TrCHSh*, *TrDFRd*, *TrF3Ha*, *TrPALa*, *TrPALb*, *TrPALf* and *TrVRa* cDNAs comprising the full open reading frame sequences in sense and antisense orientations under the control of the CaMV 35S promoter with duplicated enhancer region (35S<sup>2</sup>) were generated as detailed in Example 6.

Agrobacterium-mediated gene transfer experiments were performed using these transformation vectors.

The production of transgenic Arabidopsis plants carrying the perennial ryegrass F3OH and white clover BANa, CH1a, CH1d, CHRc, CHSa1, CHSa3, CHSc, CHSd2, CHSf, CHSh, DFRd, F3Ha, PALa, PALb, PALf and VRa cDNAs under the control of the CaMV 35S promoter with duplicated enhancer region (35S<sup>2</sup>) is described here in detail.

**Preparation of Arabidopsis plants**

Seedling punnets were filled with Debco seed raising mixture (Debco Pty. Ltd.) to form a mound. The mound was covered with two layers of anti-bird netting secured with rubber bands on each side. The soil was saturated with water and



enough seeds (*Arabidopsis thaliana* ecotype Columbia, Lehle Seeds #WT-02) sown to obtain approximately 15 plants per punnet. The seeds were then vernalised by placing the punnets at 4 °C. After 48 hours the punnets were transferred to a growth room at 22 °C under fluorescent light (constant illumination, 55  $\mu\text{molm}^{-2}\text{s}^{-1}$ ) and fed with Miracle-Gro (Scotts Australia Pty. Ltd.) once a week. Primary bolts were removed as soon as they appeared. After 4 – 6 days the secondary bolts were approximately 6 cm tall, and the plants were ready for vacuum infiltration.

### **Preparation of *Agrobacterium***

*Agrobacterium tumefaciens* strain AGL-1 were streaked on LB medium containing 50  $\mu\text{g/ml}$  rifampicin and 50  $\mu\text{g/ml}$  kanamycin and grown at 27 °C for 48 hours. A single colony was used to inoculate 5 ml of LB medium containing 50  $\mu\text{g/ml}$  rifampicin and 50  $\mu\text{g/ml}$  kanamycin and grown over night at 27 °C and 250 rpm on an orbital shaker. The overnight culture was used as an inoculum for 500 ml of LB medium containing 50  $\mu\text{g/ml}$  kanamycin only. Incubation was over night at 27 °C and 250 rpm on an orbital shaker in a 2 l Erlenmeyer flask.

The overnight cultures were centrifuged for 15 min at 5500 xg and the supernatant discarded. The cells were resuspended in 1 l of infiltration medium [5% (w/v) sucrose, 0.03% (v/v) Silwet-L77 (Vac-In-Stuff, Lehle Seeds #VIS-01)] and immediately used for infiltration.

### **Vacuum infiltration**

The *Agrobacterium* suspension was poured into a container (Décor Telfresh storer, #024) and the container placed inside the vacuum desiccator (Bel Art, #42020-0000). A punnet with *Arabidopsis* plants was inverted and dipped into the *Agrobacterium* suspension and a gentle vacuum (250 mm Hg) was applied for 2 min. After infiltration, the plants were returned to the growth room where they were kept away from direct light overnight. The next day the plants were returned to full direct light and allowed to grow until the siliques were fully developed. The plants were then allowed to dry out, the seed collected from the siliques and either

stored at room temperature in a dry container or used for selection of transformants.

### **Selection of transformants**

Prior to plating the seeds were sterilised as follows. Sufficient seeds for one  
5 150 mm petri dish (approximately 40 mg or 2000 seeds) were placed in a 1.5 ml  
microfuge tube. 500  $\mu$ l 70% ethanol were added for 2 min and replaced by 500  $\mu$ l  
sterilisation solution ( $H_2O$ :4% chlorine:5% SDS, 15:8:1). After vigorous shaking,  
the tube was left for 10 min after which time the sterilisation solution was replaced  
with 500  $\mu$ l sterile water. The tube was shaken and spun for 5 sec to sediment the  
10 seeds. The washing step was repeated 3 times and the seeds were left covered  
with approximately 200  $\mu$ l sterile water.

The seeds were then evenly spread on 150 mm petri dishes containing  
germination medium (4.61 g Murashige & Skoog salts, 10 g sucrose, 1 ml 1 M  
KOH, 2 g Phytigel, 0.5 g MES and 1 ml 1000x Gamborg's B-5 vitamins per litre)  
15 supplemented with 250  $\mu$ g/ml timetin and 75  $\mu$ g/ml gentamycin. After vernalisation  
for 48 hours at 4 °C the plants were grown under continuous fluorescent light (55  
 $\mu$ mol m<sup>-2</sup>s<sup>-1</sup>) at 22 °C to the 6 – 8 leaf stage and transferred to soil.

### **Preparation of genomic DNA**

3 – 4 leaves of Arabidopsis plants regenerated on selective medium were  
20 harvested and freeze-dried. The tissue was homogenised on a Retsch MM300  
mixer mill, then centrifuged for 10 min at 1700xg to collect cell debris. Genomic  
DNA was isolated from the supernatant using Wizard Magnetic 96 DNA Plant  
System kits (Promega) on a Biomek FX (Beckman Coulter). 5  $\mu$ l of the sample (50  
 $\mu$ l) were then analysed on an agarose gel to check the yield and the quality of the  
25 genomic DNA.

### **Analysis of DNA using real-time PCR**

Genomic DNA was analysed for the presence of the transgene by real-time  
PCR using SYBR Green chemistry. PCR primer pairs (Table 4) were designed

using MacVector (Accelrys). The forward primer was located within the 35S<sup>2</sup> promoter region and the reverse primer within the transgene to amplify products of approximately 150 - 250 bp as recommended. The positioning of the forward primer within the 35S<sup>2</sup> promoter region guaranteed that homologous genes in

5 Arabidopsis were not detected.

5  $\mu$ l of each genomic DNA sample was run in a 50  $\mu$ l PCR reaction including SYBR Green on an ABI (Applied Biosystems) together with samples containing DNA isolated from wild type Arabidopsis plants (negative control), samples containing buffer instead of DNA (buffer control) and samples containing

10 the plasmid used for transformation (positive plasmid control).

Plants were obtained after transformation with all chimeric constructs and selection on medium containing gentamycin. The selection process and two representative real-time PCR analyses are shown in Figure xx.

TABLE 4

15 **List of primers used for Real-time PCR analysis of Arabidopsis plants transformed with chimeric perennial ryegrass genes involved in flavonoid biosynthesis**

construct	primer 1 (forward)	primer 2 (reverse)
pPZP221LpF3OHsense	TTGGAGAGGACACGCTGAAATC	AGGAGAGGGTTGGACATCGC
pPZP221LpF3OHanti	CATTTTCATTTGGAGAGGACACGC	ACGAGGAGTTCTGGAAGATGGG
pPZP221TrBANasense	TTGGAGAGGACACGCTGAAATC	GCAACAAAACCAGTGCCACC
pPZP221TrBANaanti	TCATTTGGAGAGGACACGCTG	GATGATTGCCCCAGCAAGG
pPZP221TrCHlasense	CATTTTCATTTGGAGAGGACACGC	CAAGGTTCCTCGACTTGGATTGC
pPZP221TrCHlaanti	TCATTTGGAGAGGACACGCTG	AGATTACCTGCCTTGTTGAACGAG
pPZP221TrCHldsense	TCATTTGGAGAGGACACGCTG	GACGGTAGGAGGGAATAGATTGTTTC
pPZP221TrCHldanti	TCATTTGGAGAGGACACGCTG	CCAGGTATCCGAGTTATTCAACG
pPZP221TrCHRCsense	CCACTATCCTTCGCAAGACCC	TCCCATTTCCAACCACAGGC
pPZP221TrCHRCanti	TCATTTGGAGAGGACACGCTG	CAAGCCAGGACTCAGTGACCTATG
pPZP221TrCHSa1sense	TCATTTGGAGAGGACACGCTG	CTGGTCAACACGATTGCTGG
pPZP221TrCHSa1anti	TCATTTGGAGAGGACACGCTG	AACCACAGGAGAAGGACTTGACTG

pPZP221TrCHSa3sense	CATTTTCATTTGGAGAGGACACGC	AACACGGTTTGGTGGATTTC
pPZP221TrCHSa3anti	TCATTTGGAGAGGACACGCTG	ACAACCTGGAGAAGGACTTGATTGG
pPZP221TrCHScsense	TTGGAGAGGACACGCTGAAATC	ACAAGTTGGTGAGGGAATGCC
pPZP221TrCHSchanti	TCATTTGGAGAGGACACGCTG	GGGATTGATACTTGCTTTTGGACC
pPZP221TrCHSd2sense	CCCACTATCCTTCGCAAGACC	AGTTGCAGTGCCGATTGCC
pPZP221TrCHSd2anti	CATTTTCATTTGGAGAGGACACGC	AAGATGGACTTGCCACAACAGG
pPZP221TrCHSfsense	CATTTTCATTTGGAGAGGACACGC	TCGTTGCCTTTCCCTGAGTAGG
pPZP221TrCHSfanti	TCATTTGGAGAGGACACGCTG	GATTGGCTTTTGGACCAGGG
pPZP221TrCHShsense	TCATTTGGAGAGGACACGCTG	CGGTCACCATTTTTTTGTTGGAGG
pPZP221TrCHShanti	TCATTTGGAGAGGACACGCTG	TGTTGTTTGGGTTTGGACCG
pPZP221TrDFRdsense	CATTTTCATTTGGAGAGGACACGC	ATTGAGATTTTGGACGGTGGC
pPZP221TrDFRdanti	CATTTTCATTTGGAGAGGACACGC	CGCAACCTGGATTGTTGAGAGC
pPZP221TrF3Hasense	TCATTTGGAGAGGACACGCTG	TCTTCCCTAACGAACTTGACTCG
pPZP221TrF3Haanti	TCATTTGGAGAGGACACGCTG	GAACAACAACCTAGGGACTTGAGAGG
pPZP221TrPALasense	ATGACGCACAATCCCACTATCC	TTGCCTCAGCAGCCACACC
pPZP221TrPALaanti	GGAGAGGACACGCTGAAATCAC	TGCCAAAAGAGGTTGAAAGTGC
pPZP221TrPALbsense	ATCCCACTATCCTTCGCAAGACCC	AATGACTCCCCATCAACGACTCCG
pPZP221TrPALbanti	TTGGAGAGGACACGCTGAAAATC	GACAAATTGTTTACAGCTATGTGCC
pPZP221TrPALfsense	ATCCCACTATCCTTCGCAAGACCC	CACCATACGCTTCACCTCATCC
pPZP221TrPALfanti	TCATTTGGAGAGGACACGCTG	TTGTTAGAGAGGAGTTAGGAACCGC
pPZP221TrVRasense	CCACTATCCTTCGCAAGACCC	GCTTACATCCCTCTTACGTTCTGG
pPZP221TrVRaanti	CCACTATCCTTCGCAAGACCC	AAAAGCTCGTGGACGCTGG

## EXAMPLE 7

**Genetic mapping of perennial ryegrass genes involved in flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits**

The cDNAs representing genes involved in flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens;

- pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits, were amplified by PCR from their respective plasmids, gel-purified and radio-labelled for use as probes to detect restriction fragment length polymorphisms (RFLPs). RFLPs were mapped in
- 5 the F<sub>1</sub> (first generation) population, NA<sub>6</sub> x AU<sub>6</sub>. This population was made by crossing an individual (NA<sub>6</sub>) from a North African ecotype with an individual (AU<sub>6</sub>) from the cultivar Aurora, which is derived from a Swiss ecotype. Genomic DNA of the 2 parents and 114 progeny was extracted using the 1 x CTAB method of Fulton et al. (1995).
- 10 Probes were screened for their ability to detect polymorphism using the DNA (10 µg) of both parents and 5 F<sub>1</sub> progeny restricted with the enzymes DraI, EcoRI, EcoRV or HindIII. Hybridisations were carried out using the method of Sharp et al. (1988). Polymorphic probes were screened on a progeny set of 114 individuals restricted with the appropriate enzyme (Figure 115).
- 15 RFLP bands segregating within the population were scored and the data was entered into an Excel spreadsheet. Alleles showing the expected 1:1 ratio were mapped using MAPMAKER 3.0 (Lander et al. 1987). Alleles segregating from, and unique to, each parent, were mapped separately to give two different linkage maps. Markers were grouped into linkage groups at a LOD of 5.0 and
- 20 ordered within each linkage group using a LOD threshold of 2.0.
- Loci representing genes involved in flavonoid biosynthesis mapped to the linkage groups as indicated in Table 5 and in Figure 197. These gene locations can now be used as candidate genes for quantitative trait loci associated with flavonoid biosynthesis, protein binding, metal chelation, anti-oxidation, UV-light
- 25 absorption, tolerance to biotic stresses such as viruses, micro-organisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety and isoflavonoid content leading to health benefits.

TABLE 5

Map locations of ryegrass genes involved in flavonoid biosynthesis across  
two genetic linkage maps of perennial ryegrass

Probe	Polymorphic	Mapped with	Locus	Linkage group	
				NA <sub>6</sub>	AU <sub>6</sub>
<i>LpDFRb</i>	Y	<i>Hind</i> III	<i>LpDFRb</i>	6	6

5

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Finally, it is to be understood that various alterations, modifications and/or additions may be made without departing from the spirit of the present invention as outlined herein.

15 It will also be understood that the term "comprises" (or its grammatical variants) as used in this specification is equivalent to the term "includes" and should not be taken as excluding the presence of other elements or features.

Documents cited in this specification are for reference purposes only and their inclusion is not acknowledgment that they form part of the common general knowledge in the relevant art.

**CLAIMS**

1. A substantially purified or isolated nucleic acid or nucleic acid fragment encoding a flavonoid biosynthetic enzyme selected from the group consisting of chalcone isomerase (CHI), chalcone synthase (CHS), chalcone reductase (CHR), dihydroflavonol 4-reductase (DFR), leucoanthocyanidin reductase (LCR), flavonoid 3', 5' hydrolase (F3'5'H), flavanone 3-hydroxylase (F3H), flavonoid 3'-hydroxylase (F3'H), phenylalanine ammonia-lyase (PAL) and vestitone reductase (VR) from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species, or a functionally active fragment or variant thereof.

2. A nucleic acid or nucleic acid fragment according to Claim 1, wherein said clover species is white clover (*Trifolium repens*) and said ryegrass species is perennial ryegrass (*Lolium perenne*).

3. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a CHI or CHI-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 1, 3, 4, 6, 7, 9, 10, 12, 122 and 127 hereto (Sequence ID Nos: 1, 3 to 7, 8, 10 to 12, 13, 15 and 16, 17, 19 to 22, 307, and 309, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

4. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a CHS or CHS-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 13, 15, 16, 18, 19, 21, 22, 24, 25, 27, 28, 30, 31, 33, 34, 137, 142, 147, 152, 157 and 162 hereto (Sequence ID Nos: 23, 25 to 63, 64, 66 to 68, 69, 71 to 77, 78, 80 to 90, 91, 93 and 94, 95, 97 to 100, 101, 103 to 105, 106, 313, 315, 317, 319, 321, and 323, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).



5. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a CHR or CHR-like protein and including nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 36, 38, 40, 41, 43 and 132 hereto (Sequence ID Nos: 108, 110, 112 to 116, 117, 119 to 134, and  
5 311, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

6. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a DFR or DFR-like protein and including a nucleotide sequence selected  
10 from the group consisting of (a) sequences shown in Figures 44, 46, 47, 49, 50, 52, 54, 55, 57, 59, 61, 62, 64, 101, 103, 104, 106, 117 and 167 hereto (Sequence ID Nos: 135, 137 to 146, 147, 149 to 152, 153, 155, 157 and 158, 159, 161, 163, 165 to 167, 168, 170 to 184, 286, 288 to 292, 293, 295 to 297, 305, and 325, respectively); (b) complements of the sequences recited in (a); (c) sequences  
15 antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

7. A nucleic acid or nucleic acid fragment according to Claim 1, encoding an LCR or LCR-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 65 and 67  
20 hereto (Sequence ID Nos: 185 and 187 to 193, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

8. A nucleic acid or nucleic acid fragment according to Claim 1,  
25 encoding a F3'5'H or F3'5'H-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 68, 70 and 72 hereto (Sequence ID Nos: 194, 196, and 198 to 201, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants  
30 of the sequences recited in (a), (b) and (c).

9. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a F3H or F3H-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 73, 75, 76, 78, 107, 109, 111 and 172 hereto (Sequence ID Nos: 202, 204 to 244, 245, 247, 298, 300  
5 to 302, 303, and 327, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

10. A nucleic acid or nucleic acid fragment according to Claim 1,  
10 encoding a F3'H or F3'H-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 80 and 82 hereto (Sequence ID Nos: 249, and 251 and 252, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences  
15 recited in (a), (b) and (c).

11. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a PAL or PAL-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 83, 85, 86, 88, 89, 91, 93, 95, 97, 177, 182 and 187 hereto (Sequence ID Nos: 253, 255 to 257, 258,  
20 260 to 267, 268, 270, 272, 274, 276 and 277, 329, 331, and 333, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c).

12. A nucleic acid or nucleic acid fragment according to Claim 1,  
25 encoding a VR or VR-like protein and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 98, 100 and 192 hereto (Sequence ID Nos: 278, 280 to 285, and 335, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants  
30 of the sequences recited in (a), (b) and (c).

13. A construct including a nucleic acid or nucleic acid fragment according to Claim 1.

14. A vector including a nucleic acid or nucleic acid fragment according to Claim 1.

5        15. A vector according to Claim 14, further including a promoter and a terminator, said promoter, nucleic acid or nucleic acid fragment and terminator being operatively linked.

16. A plant cell, plant, plant seed or other plant part, including a construct according to claim 13 or a vector according to Claim 14.

10       17. A plant, plant seed or other plant part derived from a plant cell or plant according to Claim 16.

18. A method of modifying flavonoid biosynthesis in a plant, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment according to Claim 1, a construct according to claim 13 and/or a  
15       vector according to Claim 14.

19. A method of modifying protein binding, metal chelation, anti-oxidation, and/or UV-light absorption in a plant, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment according to Claim 1, a construct according to claim 13 and/or a vector according  
20       to Claim 14.

20. A method of modifying pigment production in a plant, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment according to Claim 1, a construct according to claim 13 and/or a vector according to Claim 14.

25       21. A method of modifying plant defense to a biotic stress, said method including introducing into said plant an effective amount of a nucleic acid or nucleic

acid fragment according to Claim 1, a construct according to claim 13 and/or a vector according to Claim 14.

22. A method according to claim 21 wherein said biotic stress is selected from the group consisting of viruses, microorganisms, insects and fungal  
5 pathogens.

23. A method of modifying forage quality of a plant by disrupting protein foam and/or conferring protection from rumen pasture bloat, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment according to Claim 1, a construct according to claim 13 and/or a vector  
10 according to Claim 14.

24. Use of a nucleic acid or nucleic acid fragment according to Claim 1, and/or nucleotide sequence information thereof, and/or single nucleotide polymorphisms thereof as a molecular genetic marker.

25. A substantially purified or isolated polypeptide from a clover  
15 (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species, selected from the group consisting of ASR and ASR-like, A22 and A22-like, CYS and CYS-like, LEA and LEA-like, DHN and DHN-like and PKABA and PKABA-like; and functionally active fragments and variants thereof.

26. A polypeptide according to Claim 25, wherein said clover species is  
20 white clover (*Trifolium repens*) and said ryegrass species is perennial ryegrass (*Lolium perenne*).

27. A polypeptide according to Claim 25, wherein said polypeptide is CHI or CHI-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 2, 5, 8, 11, 123 and 128 hereto  
25 (Sequence ID Nos: 2, 9, 14, 18, 308, and 310, respectively); and functionally active fragments and variants thereof.

28. A polypeptide according to Claim 25, wherein said polypeptide is CHS or CHS-like and includes an amino acid sequence selected from the group

consisting of sequences shown in Figures 14, 17, 20, 23, 26, 29, 32, 35, 138, 143, 148, 153, 158 and 163 hereto (Sequence ID Nos: 24, 65, 70, 79, 92, 96, 102, 107, 314, 316, 318, 320, 322, and 324, respectively); and functionally active fragments and variants thereof.

5           29. A polypeptide according to Claim 25, wherein said polypeptide is CHR or CHR-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 37, 39, 42 and 133 hereto (Sequence ID Nos: 109, 111, 118, and 312, respectively); and functionally active fragments and variants thereof.

10           30. A polypeptide according to Claim 25, wherein said polypeptide is DFR or DFR-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 45, 48, 51, 53, 56, 58, 60, 63, 102, 105, 118 and 168 hereto (Sequence ID Nos: 136, 148, 54, 156, 160, 162, 164, 169, 287, 294, 306, and 326, respectively); and functionally active fragments and  
15 variants thereof.

31. A polypeptide according to Claim 25, wherein said polypeptide is LCR or LCR-like and includes an amino acid sequence shown in Figure 66 hereto (Sequence ID No: 186); and functionally active fragments and variants thereof.

32. A polypeptide according to Claim 25, wherein said polypeptide is  
20 F3'5'H or F3'5'H-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 69 and 71 hereto (Sequence ID Nos: 195 and 197, respectively); and functionally active fragments and variants thereof.

33. A polypeptide according to Claim 25, wherein said polypeptide is  
25 F3H or F3H-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 74, 77, 79, 108, 112 and 173 hereto (Sequence ID Nos: 203, 246, 248, 299, 304, and 328, respectively); and functionally active fragments and variants thereof.

34. A polypeptide according to Claim 25, wherein said polypeptide is F3'H or F3'H-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figure 81 hereto (Sequence ID No: 250); and functionally active fragments and variants thereof.
- 5        35. A polypeptide according to Claim 25, wherein said polypeptide is PAL or PAL-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 84, 87, 90, 92, 94, 96, 178, 183 and 188 hereto (Sequence ID Nos: 254, 259, 269, 271, 273, 275, 330, 332, and 334, respectively); and functionally active fragments and variants thereof.
- 10       36. A polypeptide according to Claim 25, wherein said polypeptide is VR or VR-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 99 and 193 hereto (Sequence ID Nos: 279 and 336, respectively); and functionally active fragments and variants thereof.

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      *           20           *           40           *           60
TrCH1a : GCATTAAACANTGAAANTTGACCAGTCCCAACAAAGATCTGAAACACATAGCTCCCCATT : 60

      *           80           *           100          *           120
TrCH1a : TTTTAACATTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAA : 120

      *           140          *           160          *           180
TrCH1a : TTTCCGGCTGTGATTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG : 180

      *           200          *           220          *           240
TrCH1a : GAGAGAGGTTTGACTATTGAAGGAACTTCATCAAGTTCACTGCCATAGGAGTATATTTG : 240

      *           260          *           280          *           300
TrCH1a : GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG : 300

      *           320          *           340          *           360
TrCH1a : CTTGAGACCCTTGACTTCTACAGAGATATCATTTCAAGACCATTTGAGAAGTTGATTCGA : 360

      *           380          *           400          *           420
TrCH1a : GGATCGAAGATTAGGGAATTGAGTGGTCCTGAGTACTCAAGGAAGGTTAATGAAAACGTG : 420

      *           440          *           460          *           480
TrCH1a : GTGGCACACTTAAAATCTGTTGGGACTTATGGAGATGCAGAAGTTGAAGCTATGCAAAAA : 480

      *           500          *           520          *           540
TrCH1a : TTTGTTGAAGCCTTCAAGCCTATTAATTTTCCACCTGGTGCCTCTGTTTTTTACAGGCAA : 540

      *           560          *           580          *           600
TrCH1a : TCACCTGATGGAATATTAGGGCTTAGTTTCTCTCAAGATGCAAGTATACCAGAAAAGGAA : 600

      *           620          *
TrCH1a : GCTGCAGTAATAGAGAACAAGGGAGCTTCATCGGCG : 636

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## FIGURE 1

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TrCH1a : MSAITAIQVENLEFPVAVITSPATGKSYFLGGAGERGLTIEGNFIKFTAIGVYLEDVAVAS : 60

TrCH1a : LATKWKGKSSEELLETLDFYRDIISGPFEKLIRGSKIRELSGPEYSRKVNENCVAHLKSV : 120

TrCH1a : GTYGDAEVEAMQKFVEAFKPINFPPGASVFYRQSPDGILGLSFSQDASIPEKEAAVIENK : 180

TrCH1a : GASSA : 185

**FIGURE 2**



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		*	20	*	40	*	60		
TrCH1a1:	GCATTAAACA	TTGAGA	-NAGT	-CCNAAT	-AAAAAAGATCTGAAACACATAG	TN	CCCCATT	:	57
TrCH1a2:	-----	-----	-----	-----	-----	-----	-----	:	45
TrCH1a3:	-----	-----	-----	-----	-----	-----	-----	:	23
TrCH1a4:	-----	-----	-----	-----	-----	-----	-----	:	24
TrCH1a5:	-----	-----	-----	-----	-----	-----	-----	:	51
		*	80	*	100	*	120		
TrCH1a1:	TTTTAACATTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAA	:	117						
TrCH1a2:	TTTTAACATTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAA	:	105						
TrCH1a3:	TTTTAACATTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAA	:	83						
TrCH1a4:	TTTTAAATTTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAA	:	84						
TrCH1a5:	TTTTAACATTAAACTAAAAATATGTCGGCCATCACCGCAATCCAAGTCGAGAACCTTGAC	:	111						
		*	140	*	160	*	180		
TrCH1a1:	TTTCCGGCTGTG	TTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG	:	177					
TrCH1a2:	TTTCCGGCTGTGATTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG	:	165						
TrCH1a3:	TTTCCGGCTGTGATTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG	:	143						
TrCH1a4:	TTTCCGGCTGTGATTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG	:	144						
TrCH1a5:	TTTCCGGCTGTGATTACTTCTCCGGCCACCGGTAAGTCATATTTTCTTGGTGGTGCAGGG	:	171						
		*	200	*	220	*	240		
TrCH1a1:	GAGAGAGGTTTGACTATTGAAGGAACTTTCATCAAGTTCACTGCCATAGGAGTATATTTG	:	237						
TrCH1a2:	GAGAGAGGTTTGACTATTGAAGGAACTTTCATCAAGTTCACTGCCATAGGAGTATATTTG	:	225						
TrCH1a3:	GAGAGAGGTTTGACTATTGAAGGAACTTTCATCAAGTTCACTGCCATAGGAGTATATTTG	:	203						
TrCH1a4:	GAGAGAGGTTTGACTATTGAAGGAACTTTCATCAAGTTCACTGCCATAGGAGTATATTTG	:	204						
TrCH1a5:	GAGAGAGGTTTGACTATTGAAGGAACTTTCATCAAGTTCACTGCCATAGGAGTATATTTG	:	231						
		*	260	*	280	*	300		
TrCH1a1:	GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG	:	244						
TrCH1a2:	GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG	:	285						
TrCH1a3:	GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG	:	263						
TrCH1a4:	GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG	:	228						
TrCH1a5:	GAAGATGTAGCAGTGGCTTCACTTGCCACTAAATGGAAGGGCAAATCCTCTGAAGAGTTG	:	291						
		*	320	*	340	*	360		
TrCH1a1:	CTTGAGACCCCTTGACTTCTACAGAGATATCATTTTCAGGACCATTGAGAAGTTGATTGCA	:	-						
TrCH1a2:	CTTGAGACCCCTTGACTTCTACAGAGATATCATTTTCAGGACCATTGAGAAGTTGATTGCA	:	345						
TrCH1a3:	CTTGAGACCCCTTGACTTCTACAGAGATATCATTTTCAGGACCATTGAGAAGTTGATTGCA	:	323						
TrCH1a4:	CTTGAGACCCCTTGACTTCTACAGAGATATCATTTTCAGGACCATTGAGAAGTTGATTGCA	:	-						
TrCH1a5:	CTTGAGACCCCTTGACTTCTACAGAGATATCATTTTCAGGACCATTGAGAAGTTGATTGCA	:	351						
		*	380	*	400	*	420		
TrCH1a1:	GGATCGAAGATTAGGGAATTGAGTGGTCTGAGTACTCAAGGAAGGTTAATGAAAACCTGT	:	-						
TrCH1a2:	GGATCGAAGATTAGGGAATTGAGTGGTCTGAGTACTCAAGGAAGGTTAATGAAAACCTGT	:	405						
TrCH1a3:	GGATCGAAGATTAGGGAATTGAGTGGTCTGAGTACTCAAGGAAGGTTAATGAAAACCTGT	:	383						
TrCH1a4:	GGATCGAAGATTAGGGAATTGAGTGGTCTGAGTACTCAAGGAAGGTTAATGAAAACCTGT	:	-						
TrCH1a5:	GGATCGAAGATTAGGGAATTGAGTGGTCTGAGTACTCAAGGAAGGTTAATGAAAACCTGT	:	411						

FIGURE 3

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	*	440	*	460	*	480	
TrCH1a1:	-----						:
TrCH1a2:	GTGGCACACTTAAAAATCTGTTGGGACTTACGGAGATGCAGAAGTTGAAGCTATGCAAAAA						: 465
TrCH1a3:	GTGGCCCACTTAAAAATCTGTTGGGACTTATGGAGATGCTGAAGCTGAAGCTATGCAAAAA						: 443
TrCH1a4:	-----						:
TrCH1a5:	GTGGCACACTTAAAAATCTGTTGGGACTTATGGAGATGCAGAAGTTGAAGCTATGCAAAAA						: 471

	*	500	*	520	*	540	
TrCH1a1:	-----						:
TrCH1a2:	TTTGTTGAAGCCTTCAAGCCTATTAATTTTCACCTGGTGCCTCTGTTTTTTACAGGCAA						: 525
TrCH1a3:	TTTGTTGAAGCCTTCAAGCCTATTAATTTTCACCTGGTGCCTCTGTTTTTTACAGGCAA						: 503
TrCH1a4:	-----						:
TrCH1a5:	TTTGTTGAAGCCTTCAAGCCTATTAATTTTCACCTGGTGCCTCTGTTTTTTACAGGCAA						: 531

	*	560	*	580	*	600	
TrCH1a1:	-----						:
TrCH1a2:	TCACCTGATGGAATATTAGGGCTTAGTTTCTCTCAAGATGCAAGTATACCAGAAAAGGAA						: 585
TrCH1a3:	TCACCTGATGGAATATTAGGGCTTAGTTTCTCTCAAGATGCAAGTATACCAGAAAAGGAG						: 563
TrCH1a4:	-----						:
TrCH1a5:	TCACCTGATGGAATATTAGGGCTTAGTTTCTCTCAAGATGCAAGTATACCAGAAAAGN--						: 589

	*	620	*		
TrCH1a1:	-----		:	-	
TrCH1a2:	GCTGCAGTAATAGAGAACAANN--		:	606	
TrCH1a3:	GCTGCAGTAATAGAGAACAAGGGAGCTTCATCGGCC		:	599	
TrCH1a4:	-----		:	-	
TrCH1a5:	-----		:	-	

FIGURE 3 (cont)

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TrCH1b : TTAAAATTGACACAGTCCCAACCTTAAANTTGACNNGGTCCCAAACAAAAGATCTGAAACA : 60

TrCH1b : ACATAGCCCCCATTTTTTAACATTAAACTAAAAATATGTCTGCCATCACCGCAATCCAA : 120

TrCH1b : GTCGAGAACCTTGAATTCCCGGCGGTGATTACTTCTCCGGTCAACGGTAAGTCATATTTT : 180

TrCH1b : CTTGGTGGTGCAGGGGAGAGAGGT'TNGACTATTGAAGGAACTTCATCAAGTTCACTGCC : 240

TrCH1b : ATAGGAGTATATTTGGAAGATGTAGCAGGGGCTTCACTTGCCACTAAATGGAAGGGCAGA : 300

TrCH1b : TCCTCTGAAGAGNGCTTGAGACCC'TNGACTNC : 332

**FIGURE 4**

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```

      *           20           *           40           *           60
TrCH1b : MSAITAIQVENLEFPVITSPVNGKSYFLGGAGERGXTIEGNFIKFTAIGVYLEDVAGAS : 60

      *
TrCH1b : LATKWKGRSSEEXLRPXT : 78

```

## FIGURE 5

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```

          *           20           *           40           *           60
TrCHib1: TTAAAATTGACCNAGTCCNAACCTTAAANTTGACCNGGTCCCAAACAAAGATCTGAAACA : 60
TrCHib2: -TTAAANTTGACACAGTCCCAACCTTAAANTTGACCNGGTCCCAAACAAAGATCTGAAACA : 59
TrCHib3: -----GCGTTAAANTTGACCCAGT-CCNAACAAAGATCTGAAAC- : 38

          *           80           *           100          *           120
TrCHib1: ACATAGCCCCCATTTTTTTAACATTAAACTAAAAATATGTCTGCCATCACCGCAATCCAA : 120
TrCHib2: ACATAGCCCCCATTTTTTTAACATTAAACTAAAAATATGTCTGCCATCACCGCAATCCAA : 119
TrCHib3: ACATAGCCCCCATTTTTTTAACATTAAACTAAAAATATGTCTGCCATCACCGCAATCCAA : 98

          *           140          *           160           *           180
TrCHib1: GTCGAGAACCTTGAATTCCCGGCGGTGATTACTTCTCCGGTCAACGGTAAGTCATATTTT : 180
TrCHib2: GTCGAGAACCTTGAATTCCCGGCGGTGATTACTTCTCCGGTCAACGGTAAGTCATATTTT : 179
TrCHib3: GTCGAGAACCTTGAATTCCCGGCGGTGATTACTTCTCCGGTCAACGGTAAGTCATATTTT : 158

          *           200          *           220           *           240
TrCHib1: CTTGGTGGTGCAGGGGAGAGAGGTTTGACTATTGAAGGAAACTTCATCAAGTTCACTGCC : 240
TrCHib2: CTTGGTGGTGCAGGGGAGAGAGGTTTGACTATTGAAGGAAACTTCATCAAGTTCACTGCC : 239
TrCHib3: CTTGGTGGTGCAGGAGAGAGAGGTTTNGN----- : 186

          *           260          *           280           *           300
TrCHib1: ATAGGAGTATATTTGGAAGATGTAGCAGGGGCTTCACTTGCCACTAAATGGAAGGGGAGA : 300
TrCHib2: ATAGGAGTATATTTGGAAGATGTAGCAGGGGCTTCACTTGCCACTAAATGGAAGGGGANA : 299
TrCHib3: ----- : -

          *           320          *
TrCHib1: TCCTCTGAAGAGNCNTTGAAACCTTNGACTNN : 332
TrCHib2: TCCTCTGAAGAGGCTTGAGACCCTTGACTTC : 331
TrCHib3: ----- : -

```

FIGURE 6

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```
      *           20           *           40           *           60
TrCH1c : GTTAGNAGNAGNATNTCNNGGCACCCTTTGAAAAGTTGATTTCGAGGATCGAAGATTAGGGA : 60

      *           80           *           100          *           120
TrCH1c : ATTGAGTGGTCCTGAGTACTCAAGGAAGGTTAATGAAAAGTTCGCTGGCACACTTAAAATC : 120

      *           140          *           160          *           180
TrCH1c : TGTTGGGACTTATGGAGATGCAGAAGCTGAAGCTATGCAAAAATTTGTTGAAGCCTTCAA : 180

      *           200          *           220          *           240
TrCH1c : GCCTATTAATTTTCCACCTGGTGCCTCTGTTTTTTTACAGGCAATCACCTGATGGAATATT : 240

      *           260          *
TrCH1c : AGGGGTTAGTATTGCCAATTCATTTTTTTTAACT : 274
```

## FIGURE 7

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TrCH1c :           \*          20          \*          40          \*          60  
          : APFEKLIRGSKIRELSGPEYSRKVNENCVAHLKSVGTYGDAEAEAMQKFVEAFKPINFPP : 60

                  \*          80          \*  
TrCH1c : GASVFYRQSPDGILGVSIANSFFLTILIRVRFDC : 94

**FIGURE 8**

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	*	20	*	40	*	60	
TrCHic1:	GTTAGNAGNATNNNTTCNCGGCACCCCTTTGAAAAGTTGATTTCGAGGATCGAAGATTAGGGA						: 60
TrCHic2:	-----GNATNTTTCNCGGCACCCCTTTGAAAAGTTGATTTCGAGGATCGAAGATTAGGGA						: 50
	*	80	*	100	*	120	
TrCHic1:	ATTGAGTGGTCCTGAGTACTCAAGGAAGGTTAATGAAAAGTTCGCGTGGCACACTTAAAATC						: 120
TrCHic2:	ATTGAGTGGTCCTGAGTACTCAAGGAAGGTTAATGAAAAGTTCGCGTGGCACACTTAAAATC						: 110
	*	140	*	160	*	180	
TrCHic1:	TGTTGGGACTTATGGAGATGCAGAAGCTGAAGCTATGCAAAAATTTGTTGAAGCCTTCAA						: 180
TrCHic2:	TGTTGGGACTTATGGAGATGCAGAAGCTGAAGCTATGCAAAAATTTGTTGAAGCCTTCAA						: 170
	*	200	*	220	*	240	
TrCHic1:	GCCTATTAATTTTCCACCTGGTGCCTCTGTTTTTTTACAGGCAATCACCTGATGGAATATT						: 240
TrCHic2:	GCCTATTAATTTTCCACCTGGTGCCTCTGTTTTTTTACAGGCAATCACCTGATGGAATATT						: 230
	*	260	*				
TrCHic1:	AGGGGTTAGTATTGCCAATTCATTTTTTTTAACT						: 274
TrCHic2:	AGGGGTTAGTATTGCCAATTCATTTTTTTTAACT						: 264

FIGURE 9



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```

      *           20           *           40           *           60
TrCHId : TTNANTNNNNNTTNNCGGGCAATTACAACACACCTTCTCCATTACCATCTATCTT : 60

      *           80           *           100          *           120
TrCHId : CTACTAAGTTCAACGAGATCAATGGCACTTCCTTCTGTCAACCGCTTTGAATATCGAGAAC : 120

      *           140          *           160          *           180
TrCHId : AATCTATTCCCTCCTACCGTCACACCACCGGGATCCACCAACAATTTCTTCCTCGGCGGT : 180

      *           200          *           220          *           240
TrCHId : GCAGGAGAGCGGGGTCTTCAAATTCAAGACAAATTTGTCAAATTCACCGCTATTGGTGTT : 240

      *           260          *           280          *           300
TrCHId : TATCTACAGGACATTGCTGTTTCCTTACCTCGCCACTAAATGGAAGGGTAAGACTGCTCAA : 300

      *           320          *           340          *           360
TrCHId : GAGCTAACGGAACTGTTCCCTTTCTTCAGGGACATCGTTACAGGTCCATTTGAGAAATTT : 360

      *           380          *           400          *           420
TrCHId : ATGCAGGTGACAATGATCTTGCCATTGACTGGGCAACAATACTCAGAGAAAGTGTGAGAA : 420

      *           440          *           460          *           480
TrCHId : AATTGTGTAGCTATTTGGAAGTCTCTTGGGATTTATACCGACGAAGAAGCCAAAGCAATT : 480

      *           500          *           520          *           540
TrCHId : GAGAAGNNTGTTTCTGTCTTCAAAGANGAAACATTCCCACCAGGCTCCTCTATCCTTTTC : 540

      *           560          *           580          *           600
TrCHId : ACAGNATTACCCAAAGGATTAGGATCACTAACGATAAGNTTCTCTAAAGATGGATCCATT : 600

      *           620          *           640          *           660
TrCHId : CCAGAGACCGAGTCTGCAGTTATAGAGAATAAGCTACTCTCACAAGCTGTGCTNGAGTCG : 660

      *           680          *           700          *           720
TrCHId : ATGATAGGGGCACACGGTGTCTCCCCTGCAGCAAAACAGAGTTTTGGCCACCAGNTANC : 720

      *           740          *           760          *           780
TrCHId : CGAGNTATTCAACGAGGNTGGCTGATGCCTAGCAACTTGATNATATCAACAAAACGAAAA : 780

      *           800          *           820
TrCHId : TGAAAGNCCTTTTCTGCAATAAAGAACAAGCGGAAATTTTATTTT : 825

```

## FIGURE 10

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TrCHId : MALPSVTALNIENNLFPPPTVTPPGSTNNFFLGGAGERGLQIQDKFVKFTAIGVYLQDIAV : 60

TrCHId : PYLATKWKGKTAQELTETVPFFRDIVTGPFEKFMQVTMILPLTGQQYSEKVSENCVAIWK : 120

TrCHId : SLGIYTDEEAKAIEKVSVFKETFPPGSSILFTLPKGLGSLTIXFSKDGSI PETESAVIEN : 180

TrCHId : KLLSQAVXESMIGAHGVSPAQKQSFQGHQXXRXIQRXWLMPSNLXISTKRK : 230

## FIGURE 11

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```

      *           20           *           40           *           60
TrCHId1: TTNANTNNNNNTNNCGGCTTTNTNANAACCTACACAACACCTTCT-TTTTCCATTATCTT : 59
TrCHId2: -----GCAATTACAACCTNNCAACACCTTCTCC-TTA-CNTCTATCTT : 41
TrCHId3: -----TTAC-ACT-CACAACACCTTCTCCATTACCATCTATCTT : 37
TrCHId4: -----TGACATTATTACAATTACAACCTTAACAT : 28

      *           80           *           100          *           120
TrCHId1: CTACTAAGTTCAACGAGATCAATGGCACTTCCTTCTGTCNCCGCTTTGAATATCGAGAAC : 119
TrCHId2: CTACTAAGTTNAACGAGATCAATGGCACTTCCTTCTGTACCCGCTTTGAATATCGAGAAC : 101
TrCHId3: CTACTAAGTTCAACGAGATCAATGGCACTTCCTTCTGTACCCGCTTTGAATATCGAGAAC : 97
TrCHId4: TNACT-CTAANAAGAGATNAATGGCACTTCCTTCTGTACCCGCTTTGGATATCGAGAAC : 87

      *           140          *           160          *           180
TrCHId1: AATCTATTCCCTCCTACCGTCACACCACCGGGATCCACCAACAATTTCTTCTCGGCGGT : 179
TrCHId2: AATCTATTCCCTCCTACCGTCACACCACCGGGATCCACCAACAATTTCTTCTCGGCGGT : 161
TrCHId3: AATCTATTCCCTCCTACCGTCACACCACCGGGATCCACCAACAATTTCTTCTCGGCGGT : 157
TrCHId4: AATCTATTCCCTCCTACCGTCACACCACCGGGATCCACCAACAATTTCTTCTCGGCGGT : 147

      *           200          *           220          *           240
TrCHId1: GCAGGAGAGCGGGGTCTTCAAATTCAAGACAAATTTGTCAAATTCACCGTATTGGTGT : 239
TrCHId2: GCAGGAGAGCGGGGTCTTCAAATTCAAGACAAATTTGTCAAATTCACCGTATTGGTGT : 221
TrCHId3: GCAGGAGAGCGGGGTCTTCAAATTCAAGACAAATTTGTCAAATTCACCGTATTGGTGT : 217
TrCHId4: GCAGGAGAGCGGGGTCTTCAAATTCAAGACAAATTTGTCAAATTCACCGTATTGGTGT : 207

      *           260          *           280          *           300
TrCHId1: TATCTACAGGACATTGCTGTTCCCTTACCTCGCCACTAAATGGAAGGGTAAAGACTGCTCAA : 299
TrCHId2: TATCTACAGGACATTGCTGTTCCCTTACCTCGCCACTAAATGGAAGGGTAAAGACTGCTCAA : 266
TrCHId3: TATCTACAGGACATTGCTGTTCCCTTACCTCGCCACTAAATGGAAGGGTAAAGACTGCTCAA : 277
TrCHId4: TATCTACAGGACATTGCTGTTCCCTTACCTCGCCACTAAATGGAAGGGTAAAGACTGCTCAA : 267

      *           320          *           340          *           360
TrCHId1: GAGCTAACGGAAACTGTCCTTTCTTCAGGGACATCGTTACAGGTCCATTGAGAAATTT : 359
TrCHId2: ----- : -
TrCHId3: GAGCTAACGGAAACTGTCCTTTCTTCAGGGACATCGTTACAGGTCCATTGAGAAATTT : 337
TrCHId4: GAGCTAACGGAAACTGTCCTTTCTTCAGGGACATCGTTACAGGTCCATTGAGAAATTT : 327

      *           380          *           400          *           420
TrCHId1: ATGCAGGTGACAATGATCTTGCCATTGACTGGGCAACAATACTCAGAGAAAGTGTCAAAA : 419
TrCHId2: ----- : -
TrCHId3: ATGCAGGTGACAATGATCTTGCCATTGACTGGGCAACAATACTCAGAGAAAGTGTCAAAA : 397
TrCHId4: ATGCAGGTGACAATGATCTTGCCATTGACTGGGCAACAATACTCAGAGAAAGTGTCAAAA : 387

      *           440          *           460          *           480
TrCHId1: AATTGTGTAGCTATTTGGAAGTCTCTTGGGATTTATACCGACGAAGAAGCCAAAGCAATT : 479
TrCHId2: ----- : -
TrCHId3: AATTGTGTAGCTATTTGGAAGTCTCTTGGGATTTATACCGACGAAGAAGCCAAAGCAATT : 457
TrCHId4: AATTGTGTAGCTATTTGGAAGTCTCTTGGGATTTATACCGACGAAGAAGCCAAAGCAATT : 447

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FIGURE 12

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	*                      500                      *                      520                      *                      540	
TrCHId1:	<b>GAGAAGNNTGTTTCTGTCTTCAAAGANGAAACATTCCCACCAGGCTCCTCTATCCTTTTC</b>	: 539
TrCHId2:	-----	: -
TrCHId3:	<b>GAGAAGTTTGTTTCTGTCTTCAAAGATGAAACATTCCCACCAGGCTCCTCTATCCTTTTC</b>	: 517
TrCHId4:	<b>GANAANNNTGNTTCTGNTTNN</b> -----	: 468
	*                      560                      *                      580                      *                      600	
TrCHId1:	<b>ACAGNATTACCCAAAGGATTANGATCACTAACGATAAGNTTCTCTAAAGATGGATCCATT</b>	: 599
TrCHId2:	-----	: -
TrCHId3:	<b>ACAGNATTACCCAAAGGATTAGGATCACTAACGATAAGTTTCTCTAAAGATGGATCCATT</b>	: 577
TrCHId4:	-----	: -
	*                      620                      *                      640                      *                      660	
TrCHId1:	<b>CCAGAGACCGAGTCTGCAGTTATAGNGAATAAGCTACTCTCACAAGCTGTGCTNGAGTCG</b>	: 659
TrCHId2:	-----	: -
TrCHId3:	<b>CCAGAGACCGAGTCTGCAGTTATAGAGAATAAGCTACTCTCACAAGCTGTGCTNGAGTCG</b>	: 637
TrCHId4:	-----	: -
	*                      680                      *                      700                      *                      720	
TrCHId1:	<b>ATGATAGGGGCGANNCGGTGTCTNNCNTGCANCAAAACATAGTTTGTGNNCACCAGGNTANC</b>	: 719
TrCHId2:	-----	: -
TrCHId3:	<b>ATGATAGGGGCGCACGGTGTCTCCCCTGCAGCAAAACAGAGTTTGGCCACCAGGTTATTC</b>	: 696
TrCHId4:	-----	: -
	*                      740                      *                      760                      *                      780	
TrCHId1:	<b>CNAGNTATTCAACGAGGNTGGCTGATGCCTAGCANCTTGATNNTNTNAACAAAACNAAAA</b>	: 779
TrCHId2:	-----	: -
TrCHId3:	<b>CGAGNTATTCAACGAGGTTGGTGATGCTAGCAACTGATTATATCAACAAAACGAAAA</b>	: 753
TrCHId4:	-----	: -
	*                      800                      *                      820	
TrCHId1:	<b>TGNANGNCCTTTTCTGCAATAAAGAACA</b> -----	: 807
TrCHId2:	-----	: -
TrCHId3:	<b>TGAAAGTCCTTTCTGCAATAAAGACAAGCGGAAATTTTATTTT</b>	: 797
TrCHId4:	-----	: -

FIGURE 12 (cont)

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```

      *           20           *           40           *           60
TrCHSa: TATTNTNNGAAACCACTTGTGTTGAAGNCGTGAACCTNGCTACCCTCCATATNATACTAT : 60

      *           80           *           100          *           120
TrCHSa: NACCTCTTCTGAGACCCCTTCATCATAGAAANACAACACACNTCAGCNCTTTGCTNTTCTCT : 120

      *           140          *           160          *           180
TrCHSa: ACAACAACCTATAACTANACATATTATTTTTATNTATTTAGTATATAAATTGAAATAAACT : 180

      *           200          *           220          *           240
TrCHSa: GCTAAAGATANTTTATTAAGATATGGTGAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 240

      *           260          *           280          *           300
TrCHSa: GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTGAGCAG : 300

      *           320          *           340          *           360
TrCHSa: AGCACATATCCTGATTTCTACTTCAAAATCACAAACAGTGAGCACAAGACTGAGCTCAAA : 360

      *           380          *           400          *           420
TrCHSa: GAGAAATTCCAGCGCATGTGTGACAAATCTATGATCAAGAGCAGATACATGTATCTAACA : 420

      *           440          *           460          *           480
TrCHSa: GAAGAGATTTTGAAAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT : 480

      *           500          *           520          *           540
TrCHSa: AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT : 540

      *           560          *           580          *           600
TrCHSa: ATTAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 600

      *           620          *           640          *           660
TrCHSa: GGTGTAGACATGCCTGGTGTGATTACCAACTCACAAAACCTTAGGTCTTCGCCCATAT : 660

      *           680          *           700          *           720
TrCHSa: GTGAAGAGGTACATGATGTACCAACAAGGTGCTTTGCAGGTGGGACGGTGCTTCGTTTG : 720

      *           740          *           760          *           780
TrCHSa: GCAAAAGATTTGGCCGAGAACAACAAGGTGCTCGTGTGTTGGTTGTTTGTCTGAAGTA : 780

      *           800          *           820          *           840
TrCHSa: ACCGCAGTCACATTCCGCGGCCCCAGTGACACTCACTTGGACAGTCTTGTTGGACAAGCA : 840

      *           860          *           880          *           900
TrCHSa: CTATTTGGAGATGGAGCTGCTGCACTCATTGTTGGCTCAGACCCAGTACCAGAAATTGAG : 900

      *           920          *           940          *           960
TrCHSa: AAACCAATATTTGAGATGGTATGGACCGCACAGACAATTGCTCCAGACAGTGAAGGTGCC : 960

      *           980          *           1000         *           1020
TrCHSa: ATTGATGGTCACCTTCGTGAAGCTGGACTAACATTTTCATCTTCTTAAAGATGTTCTGCTGGG : 1020

      *           1040         *           1060         *           1080
TrCHSa: ATTGTATCAAAGAACATTAATAAAGCATTGGTCGAGGCTTTCCAACCATTAGGAATTTCT : 1080

      *           1100         *           1120         *
TrCHSa: GATTACAACCTCAATCTTTTGGATTGCACACCCGGGTGGACCTGCAATTCT : 1130

```

## FIGURE 13

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TrCHSa : MVSVAEIRKAQRAEGPATILAIGTANPANRVEQSTYPDFYFKITNSEHKTELKEKFQRM C : 60

TrCHSa : DKSMIKSRYMYLTEEILKENPSLCEYMAPSLDARQDMVVVEVPRLGKEAAVKAIKEWGQP : 120

TrCHSa : KSKITHLIFCTTSGVDMPGADYQLTKLLGLRPYVKRYMMYQQGCFAGGTVLRLAKDLAEN : 180

TrCHSa : NKGARVLVVCSEVTAVTFRGPSDTHLDSL VGQALFGDGAAALIVGSDPVPEIEKPIFEMV : 240

TrCHSa : WTAQTIAPDSEGAIDGHLREAGLTFHLLKDVPGIVSKNINKALVEAFQPLGISDYN SIFW : 300

TrCHSa : IAHPGGPAI : 309

## FIGURE 14

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	*	20	*	40	*	60	
TrCHSa1 :	TATTNTNNGAAACCACCTTGTGTTGAAGNCGTGAACCTTNGCTACCCCTCCATATNATACTAT						: 60
TrCHSa2 :	-----						: -
TrCHSa3 :	-----						: -
TrCHSa4 :	-----						: -
TrCHSa5 :	-----						: -
TrCHSa6 :	-----						: -
TrCHSa7 :	-----						: -
TrCHSa8 :	-----						: -
TrCHSa9 :	-----						: -
TrCHSa10 :	-----						: -
TrCHSa11 :	-----						: -
TrCHSa12 :	-----						: -
TrCHSa13 :	-----						: -
TrCHSa14 :	-----						: -
TrCHSa15 :	-----						: -
TrCHSa16 :	-----						: -
TrCHSa17 :	-----						: -
TrCHSa18 :	-----						: -
TrCHSa19 :	-----						: -
TrCHSa20 :	-----						: -
TrCHSa21 :	-----						: -
TrCHSa22 :	-----						: -
TrCHSa23 :	-----						: -
TrCHSa24 :	-----						: -
TrCHSa25 :	-----						: -
TrCHSa26 :	-----						: -
TrCHSa27 :	-----						: -
TrCHSa28 :	-----						: -
TrCHSa29 :	-----						: -
TrCHSa30 :	-----						: -
TrCHSa31 :	-----						: -
TrCHSa32 :	-----						: -
TrCHSa33 :	-----						: -
TrCHSa34 :	-----						: -
TrCHSa35 :	-----						: -
TrCHSa36 :	-----						: -
TrCHSa37 :	-----						: -
TrCHSa38 :	-----						: -
TrCHSa39 :	-----						: -

**FIGURE 15**

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	*	80	*	100	*	120	
TrCHSa1 :	NACCTCTTCTGAGACCCTTCATCATATATATAACNCATCTCACCTCATACATATTTCC						: 120
TrCHSa2 :	-----GNANNACAACACACATTCNAC-CTTTGCT-TTTCT						: 32
TrCHSa3 :	-----CACNTTAGCNCCTTTGCTNTTTCT						: 23
TrCHSa4 :	-----CACNTTAGC-CTTTGCT-TTTCT						: 21
TrCHSa5 :	-----GCANACACT						: 9
TrCHSa6 :	-----ACN						: 3
TrCHSa7 :	-----CC						: 2
TrCHSa8 :	-----CN						: 2
TrCHSa9 :	-----GC						: 2
TrCHSa10 :	-----						: -
TrCHSa11 :	-----						: -
TrCHSa12 :	-----						: -
TrCHSa13 :	-----						: -
TrCHSa14 :	-----						: -
TrCHSa15 :	-----						: -
TrCHSa16 :	-----						: -
TrCHSa17 :	-----						: -
TrCHSa18 :	-----						: -
TrCHSa19 :	-----						: -
TrCHSa20 :	-----						: -
TrCHSa21 :	-----						: -
TrCHSa22 :	-----						: -
TrCHSa23 :	-----						: -
TrCHSa24 :	-----						: -
TrCHSa25 :	-----						: -
TrCHSa26 :	-----						: -
TrCHSa27 :	-----						: -
TrCHSa28 :	-----						: -
TrCHSa29 :	-----						: -
TrCHSa30 :	-----						: -
TrCHSa31 :	-----						: -
TrCHSa32 :	-----						: -
TrCHSa33 :	-----						: -
TrCHSa34 :	-----						: -
TrCHSa35 :	-----						: -
TrCHSa36 :	-----						: -
TrCHSa37 :	-----						: -
TrCHSa38 :	-----						: -
TrCHSa39 :	-----						: -

FIGURE 15 (cont)



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      *           140           *           160           *           180
TrCHSa1 : ACAACAANCCTATAACTTNNCGTCTTATNTANACCAATTGAGTNTCAAATTNACATACATA :180
TrCHSa2 : ACAACAACCTTCTATNTAAACCTTTTCTTAGT-TNA--AACTTACACACTTC--CTA--AA : 85
TrCHSa3 : ACNACNNCTTCTGTCTAAACCTTTTCTGAGTNTNACTTGCATAGATAGAAANNCTANNCT : 83
TrCHSa4 : AC-AC-TCTTCTGTCTAAACNTTTTCTGAGT-TNACTTGCATACATACAAA--CTA--CT : 74
TrCHSa5 : CTATCCCTTTCTTTTCTNNAANNTAGACTACTTNGCAATAAACNACTCAATTNAATTACA : 69
TrCHSa6 : AAACAAC-TACGCATATT-ATAT-ATATA-TATAGTCTATA-ATTCAAACNAA--CT : 57
TrCHSa7 : ANGGAAACAACCTCCGNCATANTNT-TATAT-TATANNGTCTATAATTGAAAGAACT : 62
TrCHSa8 : ACAAAACACTCCCTATNCATAT-TTCTC-TCTNCTCTATA-ATTCAAAG-AAA--CT : 57
TrCHSa9 : ACAAAACACTNCGCTANTATANTATATATATNTATATATATAATTGAAAGAACT : 62
TrCHSa10 : CCNACAANCNTNAACTTNNCGTCTTATNTNNAANNATTGAGTNTNNAATTNACATACATA : 60
TrCHSa11 : AAACAAC-CCCTCATTT-ATAT-ATATA-TA-NGTCTATA-ATTCAAAG-AAA--CT : 52
TrCHSa12 : -CAACAACCN-TAACTT--CGTGTAT-TA-AAC-ATTGAGT-T-NAATT-ACATACATA : 50
TrCHSa13 : -CAACAACCTNAACTTNNCGTGTANNNTNNAACNATTGAGTNTNNAATTNACATACATA : 59
TrCHSa14 : -GANTAAACATTCCTGC-ATAT-ATAT-T-TATNTNT-TAGTCTATAATTGAAAGAACT : 58
TrCHSa15 : --AACAACCNATAACTT--CGTGTAT-TA-AAC-ATTGAGT-T-NAATT-ATATACATA : 50
TrCHSa16 : ---AAATAACCACTTNGCATATTAT-TAT-TATANNGTCTATAATTGAAAGAACT : 58
TrCHSa17 : ---NNAACCTATAACTNCC-TNTTATNT-TA-NCATTGAGTNT-NAATTNATATACATA : 57
TrCHSa18 : ----GGNTAAACAACCNLANAGTNC-TTATNTNTTNNATNTNNAAT-ATATACATA : 56
TrCHSa19 : ----TACAAAACACNCCGNATATNT-TATNTNTTGT-TATAATTGAAAGAACT : 55
TrCHSa20 : -----TACATNCGCNATAT-TAT-T-TAT-TNT-TGT-TATAATTGAAAGAACT : 53
TrCHSa21 : -----TATAACCTNCCGA-TATNT-TANNNT-T-NGT-TATAATTGAAAGAACT : 52
TrCHSa22 : -----TAAACNCNCG-NTTATNTTNTNTNTNT-TGT-TATAATTGAAAGAACT : 49
TrCHSa23 : -----CCNATN-ACCTTC-TTATNTTAC-TATAC-TACATACATA-CA : 45
TrCHSa24 : -----ACNNANACNTA-TTATNTTAC-TATAC-TACATAGCAGACT : 47
TrCHSa25 : -----ACNCNCGCTATNT-TTAT-TATNTTACT-TATAATTGAAAGAACT : 47
TrCHSa26 : -----AC-TA-NAC-TA-TTT-T-ATTACTNTACT-CAATAG-NGGACT : 40
TrCHSa27 : -----ACCTACCTTC-TTAT-NTT--N-A-ATT-CATATNG-CA : 37
TrCHSa28 : -----ACNACTTC-TTATCACTTNC-ANATACATATATAC : 42
TrCHSa29 : -----TNTNGTCTATTG-AAGAACT : 21
TrCHSa30 : -----GN : 2
TrCHSa31 : -----GT : 2
TrCHSa32 : ----- : -
TrCHSa33 : ----- : -
TrCHSa34 : ----- : -
TrCHSa35 : -----AAG : 3
TrCHSa36 : -----G : 1
TrCHSa37 : ----- : -
TrCHSa38 : ----- : -
TrCHSa39 : ----- : -

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FIGURE 15 (cont)

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      *           200           *           220           *           240
TrCHSa1 : GCGGAACATAGTAAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 240
TrCHSa2 : G--ATATA----TATCAAC--ATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 137
TrCHSa3 : GCTAAAGANNNTATCAAGNATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 143
TrCHSa4 : GCTAAAGA----TATCAAG--ATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 128
TrCHSa5 : TAAAAATCTTACTATTAAGATATGGTGAAGTGTATCTGAAATTCGTAAGGCTCAAAGGGCT : 129
TrCHSa6 : GCTAAAGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 116
TrCHSa7 : ACTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 122
TrCHSa8 : GCTAAAGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 116
TrCHSa9 : ACTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 122
TrCHSa10 : GCGGAACATAGTAAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 120
TrCHSa11 : GCTAAAGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 111
TrCHSa12 : GCGGAACATAGTAAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 110
TrCHSa13 : GCGGAACNTATTAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 119
TrCHSa14 : GCTAAAGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 117
TrCHSa15 : GCGGAACATAGTAAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 110
TrCHSa16 : ACTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 118
TrCHSa17 : GCGGAACATAGTAAATACAAGATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 117
TrCHSa18 : GCNAAA--ATAG--TATTAAGATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 116
TrCHSa19 : GCTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 115
TrCHSa20 : GCTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 113
TrCHSa21 : GCTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 112
TrCHSa22 : NCTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 109
TrCHSa23 : G--CAGCAAGATATTAAGATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 103
TrCHSa24 : GCAAAATCAAGTATTAAGATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 107
TrCHSa25 : GCTAAAGATANTTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 107
TrCHSa26 : GCAAG--TATNAGTAT--AAATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 98
TrCHSa27 : GC--AAACAAAG--TAT--AAATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 95
TrCHSa28 : GCNAAA--CAAG--TAT--AAATATGGTTAGTGTCTCTGAAATTCGCAAGGCTCAAAGGGCT : 102
TrCHSa29 : GCTA----AGA--TTT--TTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 77
TrCHSa30 : GCTAAAGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 61
TrCHSa31 : NCTAAA--NNN--TTT--TTAAGATATGGTGAAGTGTAGCTG--AATTCGC--AGGCTCAGAGGGCT : 58
TrCHSa32 : GATA--AGATA--TTATTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 58
TrCHSa33 : --TAAAGATANN--TTT--TTAAGATATGGTGAAGTGTAGCTGAAATTCGCAAGGCTCAGAGGGCT : 58
TrCHSa34 : GGNTCANA-----GGGCT : 13
TrCHSa35 : GCCC----- : 11
TrCHSa36 : GCCC----- : 9
TrCHSa37 : ---A--AAG----- : 5
TrCHSa38 : ---AT--A----- : 4
TrCHSa39 : ---GCT-- : 4

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FIGURE 15 (cont)

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      *           260           *           280           *           300
TrCHSa1 : GAAGGCCCTGCAACATATTTGGCCATTGGCTACTGCAAATCCAGCAAATCGTGTTCAG :300
TrCHSa2 : GAAGGCCCTGCAACTATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :197
TrCHSa3 : GAAGGCCCTGCAACAATCTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :203
TrCHSa4 : GAAGGCCCTGCAACAATCTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :188
TrCHSa5 : GAAGGCCCTGCAACCATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :189
TrCHSa6 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :176
TrCHSa7 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :182
TrCHSa8 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :176
TrCHSa9 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :182
TrCHSa10 : GAAGGCCCTGCAACATATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :180
TrCHSa11 : GAAGGCCCTGCAACCATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :171
TrCHSa12 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAATCGTGTTCAG :170
TrCHSa13 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAATCGTGTTCAG :179
TrCHSa14 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :177
TrCHSa15 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAATCGTGTTCAG :170
TrCHSa16 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :178
TrCHSa17 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAATCGTGTTCAG :177
TrCHSa18 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :176
TrCHSa19 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :175
TrCHSa20 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :173
TrCHSa21 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :172
TrCHSa22 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :169
TrCHSa23 : GAAGGCCCTGCAACCATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :163
TrCHSa24 : GAAGGCCCTGCAACCATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :167
TrCHSa25 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :167
TrCHSa26 : GAAGGCCCTGCAACCATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :158
TrCHSa27 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :155
TrCHSa28 : GAAGGCCCTGCAACATATTTTGGCCATTGGCTACTGCAAATCCAGCAAACCGTGTTCAG :162
TrCHSa29 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :137
TrCHSa30 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :121
TrCHSa31 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :118
TrCHSa32 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :118
TrCHSa33 : GAAGGCCCTGCAACCATTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG :118
TrCHSa34 : GAAGGCCCTGCAACC-TTTTGGCCATTGGCACTGCAAATCCAGCAAACCGTGTTCAG : 71
TrCHSa35 : -----AATTTTGGCTTTGGCTACTGCAAATCCAGCAAATCGTGTTCAG : 59
TrCHSa36 : -----ACCTTTTGGCTTTGGCTACTGCAAATCCAGCAAATCGTGTTCAG : 57
TrCHSa37 : ----- : -
TrCHSa38 : ----- : -
TrCHSa39 : ----- : -

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FIGURE 15 (cont)

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      *           320           *           340           *           360
TrCHSa1 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 360
TrCHSa2 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 257
TrCHSa3 : AGCACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 263
TrCHSa4 : AGCACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 248
TrCHSa5 : AGCACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 249
TrCHSa6 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 236
TrCHSa7 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 242
TrCHSa8 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 236
TrCHSa9 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 242
TrCHSa10 : AGCACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 240
TrCHSa11 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 231
TrCHSa12 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 230
TrCHSa13 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 239
TrCHSa14 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 237
TrCHSa15 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 230
TrCHSa16 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 238
TrCHSa17 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 237
TrCHSa18 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 236
TrCHSa19 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 235
TrCHSa20 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 233
TrCHSa21 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 232
TrCHSa22 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 229
TrCHSa23 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 223
TrCHSa24 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 227
TrCHSa25 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 227
TrCHSa26 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 218
TrCHSa27 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 215
TrCHSa28 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTCAAA : 222
TrCHSa29 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 197
TrCHSa30 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 181
TrCHSa31 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 178
TrCHSa32 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 178
TrCHSa33 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 178
TrCHSa34 : AGCACATATCCTGATTTCTACTTCAAAATACAAACAGTGAGCACAAGACTGAGCTCAAA : 131
TrCHSa35 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 119
TrCHSa36 : AGTACATATCCTGATTTCTACTTCAAAATCACTAACAGTGAGCATAAAGCTTGAGCTTAAA : 117
TrCHSa37 : -----CTGAGCTTAAA : 17
TrCHSa38 : ----- : -
TrCHSa39 : ----- : -

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FIGURE 15 (cont)

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	*	380	*	400	*	420	
TrCHSa1 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 420
TrCHSa2 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 317
TrCHSa3 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 323
TrCHSa4 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 308
TrCHSa5 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 309
TrCHSa6 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 296
TrCHSa7 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 302
TrCHSa8 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 296
TrCHSa9 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 302
TrCHSa10 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 300
TrCHSa11 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 291
TrCHSa12 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 290
TrCHSa13 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 299
TrCHSa14 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 297
TrCHSa15 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 290
TrCHSa16 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 298
TrCHSa17 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 297
TrCHSa18 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 296
TrCHSa19 :	GAGAAATTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 295
TrCHSa20 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 293
TrCHSa21 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 292
TrCHSa22 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 289
TrCHSa23 :	GAGAAATTCCAGCGCATGTGTGACAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 283
TrCHSa24 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 287
TrCHSa25 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 287
TrCHSa26 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 278
TrCHSa27 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 275
TrCHSa28 :	GAGAAATTCCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 282
TrCHSa29 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 257
TrCHSa30 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 241
TrCHSa31 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 238
TrCHSa32 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 238
TrCHSa33 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 238
TrCHSa34 :	GAGAAGTTCCACCGCATGTGTGACAAATCCATGATCAAGAGCAGATACATGTATCTAACA						: 191
TrCHSa35 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 179
TrCHSa36 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 177
TrCHSa37 :	GAGAAATTTCAGCGCATGTGTGATAAATCTATGATCAAGAGCAGATACATGTATCTAACA						: 77
TrCHSa38 :	-----AGAGCAGATACATGTATCTAACA						: 27
TrCHSa39 :	-----						: -

FIGURE 15 (cont)

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	*	440	*	460	*	480	
TrCHSa1 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 480
TrCHSa2 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 377
TrCHSa3 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAGCACATGGCACCTTCATTGGATGCT						: 383
TrCHSa4 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAGCACATGGCACCTTCATTGGATGCT						: 368
TrCHSa5 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 369
TrCHSa6 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 356
TrCHSa7 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 362
TrCHSa8 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 356
TrCHSa9 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 362
TrCHSa10 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 360
TrCHSa11 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 351
TrCHSa12 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 350
TrCHSa13 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 359
TrCHSa14 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 357
TrCHSa15 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 350
TrCHSa16 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 358
TrCHSa17 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 357
TrCHSa18 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 356
TrCHSa19 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 355
TrCHSa20 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 353
TrCHSa21 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 352
TrCHSa22 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 349
TrCHSa23 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 343
TrCHSa24 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 347
TrCHSa25 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 347
TrCHSa26 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 338
TrCHSa27 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 335
TrCHSa28 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 342
TrCHSa29 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 317
TrCHSa30 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 301
TrCHSa31 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 298
TrCHSa32 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 298
TrCHSa33 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 298
TrCHSa34 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 251
TrCHSa35 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 239
TrCHSa36 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 237
TrCHSa37 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 137
TrCHSa38 :	GAAGAGATTTTGAAGAGAAAATCCTAGTCTTTGTGAATACATGGCACCTTCATTGGATGCT						: 87
TrCHSa39 :	-----						:

FIGURE 15 (cont)



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	*	500	*	520	*	540	
TrCHSa1 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 540
TrCHSa2 :	AGGCAAGATATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 437
TrCHSa3 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 443
TrCHSa4 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 428
TrCHSa5 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTAGGGAAGGAGGCTGCAGTGAAGGCC						: 429
TrCHSa6 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCC						: 416
TrCHSa7 :	AGNCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 407
TrCHSa8 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 416
TrCHSa9 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 422
TrCHSa10 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 420
TrCHSa11 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 411
TrCHSa12 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 410
TrCHSa13 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 419
TrCHSa14 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCC						: 417
TrCHSa15 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 390
TrCHSa16 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 418
TrCHSa17 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 417
TrCHSa18 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 416
TrCHSa19 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 415
TrCHSa20 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 413
TrCHSa21 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 412
TrCHSa22 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 409
TrCHSa23 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 403
TrCHSa24 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 407
TrCHSa25 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 407
TrCHSa26 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 398
TrCHSa27 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 395
TrCHSa28 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 402
TrCHSa29 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 377
TrCHSa30 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 361
TrCHSa31 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 358
TrCHSa32 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 358
TrCHSa33 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 358
TrCHSa34 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 311
TrCHSa35 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 299
TrCHSa36 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 297
TrCHSa37 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 197
TrCHSa38 :	AGGCAAGACATGGTGGTGGTTGAGGTACCTAGACTTGGGAAGGAGGCTGCAGTGAAGGCT						: 147
TrCHSa39 :	-----						: -

FIGURE 15 (cont)

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      *           560           *           580           *           600
TrCHSa1 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 600
TrCHSa2 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 497
TrCHSa3 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 503
TrCHSa4 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 488
TrCHSa5 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 489
TrCHSa6 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 476
TrCHSa7 : ----- : -
TrCHSa8 : ATCAAAGAATGGGGTCAACCAAAGTCAAGATNACTNACTTAATCTTTTGCACCACAAGN : 476
TrCHSa9 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 482
TrCHSa10 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 480
TrCHSa11 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 471
TrCHSa12 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 470
TrCHSa13 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 479
TrCHSa14 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 477
TrCHSa15 : ----- : -
TrCHSa16 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 478
TrCHSa17 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 477
TrCHSa18 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 437
TrCHSa19 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 475
TrCHSa20 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 473
TrCHSa21 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 472
TrCHSa22 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 469
TrCHSa23 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 463
TrCHSa24 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 467
TrCHSa25 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 467
TrCHSa26 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 458
TrCHSa27 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 455
TrCHSa28 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 462
TrCHSa29 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 437
TrCHSa30 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 421
TrCHSa31 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 418
TrCHSa32 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 418
TrCHSa33 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 418
TrCHSa34 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 371
TrCHSa35 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 359
TrCHSa36 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 357
TrCHSa37 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 257
TrCHSa38 : ATCAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 207
TrCHSa39 : ---CAAAGAATGGGGTCAACCAAAGTCAAAGATTACTCACTTAATCTTTTGCACCACAAGT : 61

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FIGURE 15 (cont)



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	*	620	*	640	*	660	
TrCHSa1 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 660
TrCHSa2 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAAGCCACA	AAACTCTTAGGACTTCG	TCCATAT		: 557
TrCHSa3 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 563
TrCHSa4 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 548
TrCHSa5 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTGGACTTCG	TCCATAT		: 549
TrCHSa6 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 536
TrCHSa7 :	-----						: -
TrCHSa8 :	GGNGCCN	-----					: 483
TrCHSa9 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGACTTCG	TCCATAT		: 542
TrCHSa10 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 540
TrCHSa11 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 531
TrCHSa12 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 530
TrCHSa13 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 539
TrCHSa14 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 537
TrCHSa15 :	-----						: -
TrCHSa16 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGACTTCG	CCCATAT		: 538
TrCHSa17 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 537
TrCHSa18 :	-----						: -
TrCHSa19 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 535
TrCHSa20 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 533
TrCHSa21 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTGGACTTCG	TCCATAT		: 532
TrCHSa22 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 529
TrCHSa23 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 523
TrCHSa24 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 527
TrCHSa25 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 527
TrCHSa26 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 518
TrCHSa27 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 515
TrCHSa28 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 522
TrCHSa29 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 497
TrCHSa30 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 481
TrCHSa31 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 478
TrCHSa32 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 478
TrCHSa33 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 478
TrCHSa34 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 431
TrCHSa35 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 419
TrCHSa36 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 417
TrCHSa37 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 317
TrCHSa38 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 267
TrCHSa39 :	GGTGT	TGACATGCCTGGT	GCTGATTACCAACTCACA	AAACTCTTAGGTCTTCG	CCCATAT		: 121

FIGURE 15 (cont)

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          *           680           *           700           *           720
TrCHSa1 : GTG-----: 663
TrCHSa2 : GTGAAGAGGTACATGATG-----: 575
TrCHSa3 : GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGG-----: 607
TrCHSa4 : GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTT--: 606
TrCHSa5 : GTGAAGAGGTTTCATGATGTACCAACAAGGTTGTTTTGCAGGAGGC-----: 594
TrCHSa6 : GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 596
TrCHSa7 : -----: -
TrCHSa8 : -----: -
TrCHSa9 : GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGG-----: 586
TrCHSa10: GTGAAGAGGTACATGATGTACCAACAAGGGTGCT-----: 574
TrCHSa11: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 591
TrCHSa12: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAG-----: 570
TrCHSa13: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTT--: 598
TrCHSa14: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 597
TrCHSa15: -----: -
TrCHSa16: GTGAAGAGGCGCGTGTGNNCCN-----: 561
TrCHSa17: GTGAAG-----: 543
TrCHSa18: -----: -
TrCHSa19: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 595
TrCHSa20: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACG-----: 581
TrCHSa21: GTGAAGAGGTTCATGATGTACCAACAAGGTTGCTTTGCAGGTGGGACGGTTCCTCG-----: 588
TrCHSa22: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 589
TrCHSa23: GTGAAGAGGTATATGATGTAT-----: 544
TrCHSa24: GTGAAGAGGTACATGATGTACCAACAAG-----: 555
TrCHSa25: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAG-----: 570
TrCHSa26: GTGAAGAGGTACATGATGTACCAACAAG-----: 546
TrCHSa27: GTGAAGAGGTATATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTGCTTCGTTTG: 575
TrCHSa28: GTGAAGAGGTATATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTGCTTCNTTTG: 582
TrCHSa29: GTGAAGAGGTATATGATGTACCAA-----: 521
TrCHSa30: GTGAAGAGGTATATGATGTACCAACN-----: 506
TrCHSa31: GTGAAGAGGTATATGATGTACCAACN-----: 504
TrCHSa32: GTGAAGAGGTATATGATGTACCAACAAGGTTGTTTTGCAGGAGGCACGGTGCTTCGTTTG: 538
TrCHSa33: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 538
TrCHSa34: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 491
TrCHSa35: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 479
TrCHSa36: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 477
TrCHSa37: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 377
TrCHSa38: GTGAAGAGGTACATGATGTACCAACAAGGGTGCTTTGCAGGTGGGACGGTTCCTCGTTTG: 327
TrCHSa39: GTGAAGAGGTATATGATGTACCAACAAGGTTGCTTTGCAGGAGGCACGGTGCTTCGTTTG: 181

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FIGURE 15 (cont)

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	*	740	*	760	*	780														
TrCHSa1 :	-----		-----		-----		-													
TrCHSa2 :	-----		-----		-----		-													
TrCHSa3 :	-----		-----		-----		-													
TrCHSa4 :	-----		-----		-----		-													
TrCHSa5 :	-----		-----		-----		-													
TrCHSa6 :	GC	AAAA	GATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	GCT	AG	TTG	TTG	TCT	GAA	GTC	: 656	
TrCHSa7 :	-----		-----		-----		-													
TrCHSa8 :	-----		-----		-----		-													
TrCHSa9 :	-----		-----		-----		-													
TrCHSa10 :	-----		-----		-----		-													
TrCHSa11 :	GC	AAAA	GATT	TGG	-----		: 604													
TrCHSa12 :	-----		-----		-----		-													
TrCHSa13 :	-----		-----		-----		-													
TrCHSa14 :	GC	AAAA	GATT	TGG	-----		: 609													
TrCHSa15 :	-----		-----		-----		-													
TrCHSa16 :	-----		-----		-----		-													
TrCHSa17 :	-----		-----		-----		-													
TrCHSa18 :	-----		-----		-----		-													
TrCHSa19 :	GC	AAAA	GATT	TGG	-----		: 607													
TrCHSa20 :	-----		-----		-----		-													
TrCHSa21 :	-----		-----		-----		-													
TrCHSa22 :	GC	AAAA	GATT	TGG	CCG	AAGA	CAACA	-----		: 613										
TrCHSa23 :	-----		-----		-----		-													
TrCHSa24 :	-----		-----		-----		-													
TrCHSa25 :	-----		-----		-----		-													
TrCHSa26 :	-----		-----		-----		-													
TrCHSa27 :	GC	AA	AGC	-----			: 582													
TrCHSa28 :	GC	AA	AGC	GATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	NC	NG	TTG	CG	TTG	CT	NA	ATC	: 642
TrCHSa29 :	-----		-----		-----		-													
TrCHSa30 :	-----		-----		-----		-													
TrCHSa31 :	-----		-----		-----		-													
TrCHSa32 :	GC	AAAA	GATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 598	
TrCHSa33 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 597	
TrCHSa34 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 551	
TrCHSa35 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 539	
TrCHSa36 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 537	
TrCHSa37 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 437	
TrCHSa38 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 387	
TrCHSa39 :	GC	TAA	AGATT	TGG	CCG	AAGA	CAACA	AAAG	GTG	CTCG	TG	TG	TTG	TTG	TTG	TCT	GAA	GTC	: 241	

FIGURE 15 (cont)

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	*	800	*	820	*	840	
TrCHSa1 :	-----		-----		-----		-
TrCHSa2 :	-----		-----		-----		-
TrCHSa3 :	-----		-----		-----		-
TrCHSa4 :	-----		-----		-----		-
TrCHSa5 :	-----		-----		-----		-
TrCHSa6 :	ACCGCAGTCACATTTCGCGGGCCCCAGTGATACTCACTTGGACAGTCTTCGNTG						: 708
TrCHSa7 :	-----		-----		-----		-
TrCHSa8 :	-----		-----		-----		-
TrCHSa9 :	-----		-----		-----		-
TrCHSa10 :	-----		-----		-----		-
TrCHSa11 :	-----		-----		-----		-
TrCHSa12 :	-----		-----		-----		-
TrCHSa13 :	-----		-----		-----		-
TrCHSa14 :	-----		-----		-----		-
TrCHSa15 :	-----		-----		-----		-
TrCHSa16 :	-----		-----		-----		-
TrCHSa17 :	-----		-----		-----		-
TrCHSa18 :	-----		-----		-----		-
TrCHSa19 :	-----		-----		-----		-
TrCHSa20 :	-----		-----		-----		-
TrCHSa21 :	-----		-----		-----		-
TrCHSa22 :	-----		-----		-----		-
TrCHSa23 :	-----		-----		-----		-
TrCHSa24 :	-----		-----		-----		-
TrCHSa25 :	-----		-----		-----		-
TrCHSa26 :	-----		-----		-----		-
TrCHSa27 :	-----		-----		-----		-
TrCHSa28 :	ACCGCAN		-----		-----		: 649
TrCHSa29 :	-----		-----		-----		-
TrCHSa30 :	-----		-----		-----		-
TrCHSa31 :	-----		-----		-----		-
TrCHSa32 :	ACGCGAGTTCACATTCCGTCGGCCCCAGTGACACTCACTTGGACAGTCTTGTTGGACAAGCA						: 658
TrCHSa33 :	-----		-----		-----		-
TrCHSa34 :	ACGCGAGTCACATTCCGTCGGCCCCAGTGACACTCACTTG						: 590
TrCHSa35 :	ACCGCAGTCACATTCCGCGGGCCCCAGTGACACTCACTTGGACAGCTTGTTGGACAAGCA						: 599
TrCHSa36 :	ACCGCAGTCACATTCCGCGGGCCCCAGTGACACTCACTTGGACAGCTTGTTGGACAAGCA						: 597
TrCHSa37 :	ACCGCAGTCACATTCCGCGGGCCCCAGTGACACTCACTTGGACAGTCTTGTTGGACAAGCA						: 497
TrCHSa38 :	ACCGCAGTCACATTCCGTCGGCCCCAGTGACACTCACTTGGACAGTCTTGTTGGACAAGCA						: 447
TrCHSa39 :	ACGCGAGTCACATTCCGCGGGCCCCAGTGACACTCACTTGGACAGTCTTGTTGGACAAGCA						: 301

FIGURE 15 (cont)

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	*	860	*	880	*	900	
TrCHSa1 :	-----						:
TrCHSa2 :	-----						:
TrCHSa3 :	-----						:
TrCHSa4 :	-----						:
TrCHSa5 :	-----						:
TrCHSa6 :	-----						:
TrCHSa7 :	-----						:
TrCHSa8 :	-----						:
TrCHSa9 :	-----						:
TrCHSa10 :	-----						:
TrCHSa11 :	-----						:
TrCHSa12 :	-----						:
TrCHSa13 :	-----						:
TrCHSa14 :	-----						:
TrCHSa15 :	-----						:
TrCHSa16 :	-----						:
TrCHSa17 :	-----						:
TrCHSa18 :	-----						:
TrCHSa19 :	-----						:
TrCHSa20 :	-----						:
TrCHSa21 :	-----						:
TrCHSa22 :	-----						:
TrCHSa23 :	-----						:
TrCHSa24 :	-----						:
TrCHSa25 :	-----						:
TrCHSa26 :	-----						:
TrCHSa27 :	-----						:
TrCHSa28 :	-----						:
TrCHSa29 :	-----						:
TrCHSa30 :	-----						:
TrCHSa31 :	-----						:
TrCHSa32 :	CTATTGGAGATGGAGCTGCTGCACTCATCGTTGGCTCAGACCAGTACCAGAAATTGAG						: 718
TrCHSa33 :	-----						:
TrCHSa34 :	-----						:
TrCHSa35 :	CTATTGGAGATGGAGCTG-----						: 618
TrCHSa36 :	CTATTGGAGATGGAGCTGCTG-----						: 619
TrCHSa37 :	CTATTGGAGATGGAGCTGCTGCACTCATCGTTGGCTCAGACCAGTACCAGAAATTGAG						: 557
TrCHSa38 :	CTATTGGAGATGGAGCTGCTGCTCATCGTTGGCTCAGACCAGTACCAGAAATTGAG						: 507
TrCHSa39 :	CTATTGGAGATGGAGCTGCTGCTCATCGTTGGCTCAGACCAGTACCAGAAATTGAG						: 361

FIGURE 15 (cont)

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	*	920	*	940	*	960	
TrCHSa1 :	-----		-----		-----		:
TrCHSa2 :	-----		-----		-----		:
TrCHSa3 :	-----		-----		-----		:
TrCHSa4 :	-----		-----		-----		:
TrCHSa5 :	-----		-----		-----		:
TrCHSa6 :	-----		-----		-----		:
TrCHSa7 :	-----		-----		-----		:
TrCHSa8 :	-----		-----		-----		:
TrCHSa9 :	-----		-----		-----		:
TrCHSa10 :	-----		-----		-----		:
TrCHSa11 :	-----		-----		-----		:
TrCHSa12 :	-----		-----		-----		:
TrCHSa13 :	-----		-----		-----		:
TrCHSa14 :	-----		-----		-----		:
TrCHSa15 :	-----		-----		-----		:
TrCHSa16 :	-----		-----		-----		:
TrCHSa17 :	-----		-----		-----		:
TrCHSa18 :	-----		-----		-----		:
TrCHSa19 :	-----		-----		-----		:
TrCHSa20 :	-----		-----		-----		:
TrCHSa21 :	-----		-----		-----		:
TrCHSa22 :	-----		-----		-----		:
TrCHSa23 :	-----		-----		-----		:
TrCHSa24 :	-----		-----		-----		:
TrCHSa25 :	-----		-----		-----		:
TrCHSa26 :	-----		-----		-----		:
TrCHSa27 :	-----		-----		-----		:
TrCHSa28 :	-----		-----		-----		:
TrCHSa29 :	-----		-----		-----		:
TrCHSa30 :	-----		-----		-----		:
TrCHSa31 :	-----		-----		-----		:
TrCHSa32 :	AAACCAATATTTGAGATGGTATGGACATGCACAAACAATTGCTCCAGACAGTGAAGGTGCC						: 778
TrCHSa33 :	-----		-----		-----		:
TrCHSa34 :	-----		-----		-----		:
TrCHSa35 :	-----		-----		-----		:
TrCHSa36 :	-----		-----		-----		:
TrCHSa37 :	AN-----		-----		-----		: 559
TrCHSa38 :	AAGCCAATATTTGAGATGGTATGGACCGCACAGACAATTGCTCCAG-----						: 553
TrCHSa39 :	AAACCAATATTTGAGATGGTATGGACCGCACAGACAATTGCTCCAGACAGTGAAGGTGCC						: 421

FIGURE 15 (cont)

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	*	980	*	1000	*	1020	
TrCHSa1 :	-----		-----		-----		:
TrCHSa2 :	-----		-----		-----		:
TrCHSa3 :	-----		-----		-----		:
TrCHSa4 :	-----		-----		-----		:
TrCHSa5 :	-----		-----		-----		:
TrCHSa6 :	-----		-----		-----		:
TrCHSa7 :	-----		-----		-----		:
TrCHSa8 :	-----		-----		-----		:
TrCHSa9 :	-----		-----		-----		:
TrCHSa10 :	-----		-----		-----		:
TrCHSa11 :	-----		-----		-----		:
TrCHSa12 :	-----		-----		-----		:
TrCHSa13 :	-----		-----		-----		:
TrCHSa14 :	-----		-----		-----		:
TrCHSa15 :	-----		-----		-----		:
TrCHSa16 :	-----		-----		-----		:
TrCHSa17 :	-----		-----		-----		:
TrCHSa18 :	-----		-----		-----		:
TrCHSa19 :	-----		-----		-----		:
TrCHSa20 :	-----		-----		-----		:
TrCHSa21 :	-----		-----		-----		:
TrCHSa22 :	-----		-----		-----		:
TrCHSa23 :	-----		-----		-----		:
TrCHSa24 :	-----		-----		-----		:
TrCHSa25 :	-----		-----		-----		:
TrCHSa26 :	-----		-----		-----		:
TrCHSa27 :	-----		-----		-----		:
TrCHSa28 :	-----		-----		-----		:
TrCHSa29 :	-----		-----		-----		:
TrCHSa30 :	-----		-----		-----		:
TrCHSa31 :	-----		-----		-----		:
TrCHSa32 :	ATTG		-----		-----		: 782
TrCHSa33 :	-----		-----		-----		:
TrCHSa34 :	-----		-----		-----		:
TrCHSa35 :	-----		-----		-----		:
TrCHSa36 :	-----		-----		-----		:
TrCHSa37 :	-----		-----		-----		:
TrCHSa38 :	-----		-----		-----		:
TrCHSa39 :	ATTGATGGTCACCTTCGTGAAGCTGGACTAACATTTTCATCTTCTTAAAGATGTTCTCTGGG						: 481

## FIGURE 15 (cont)

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	*	1040	*	1060	*	1080	
TrCHSa1 :	-----		-----		-----		:
TrCHSa2 :	-----		-----		-----		:
TrCHSa3 :	-----		-----		-----		:
TrCHSa4 :	-----		-----		-----		:
TrCHSa5 :	-----		-----		-----		:
TrCHSa6 :	-----		-----		-----		:
TrCHSa7 :	-----		-----		-----		:
TrCHSa8 :	-----		-----		-----		:
TrCHSa9 :	-----		-----		-----		:
TrCHSa10 :	-----		-----		-----		:
TrCHSa11 :	-----		-----		-----		:
TrCHSa12 :	-----		-----		-----		:
TrCHSa13 :	-----		-----		-----		:
TrCHSa14 :	-----		-----		-----		:
TrCHSa15 :	-----		-----		-----		:
TrCHSa16 :	-----		-----		-----		:
TrCHSa17 :	-----		-----		-----		:
TrCHSa18 :	-----		-----		-----		:
TrCHSa19 :	-----		-----		-----		:
TrCHSa20 :	-----		-----		-----		:
TrCHSa21 :	-----		-----		-----		:
TrCHSa22 :	-----		-----		-----		:
TrCHSa23 :	-----		-----		-----		:
TrCHSa24 :	-----		-----		-----		:
TrCHSa25 :	-----		-----		-----		:
TrCHSa26 :	-----		-----		-----		:
TrCHSa27 :	-----		-----		-----		:
TrCHSa28 :	-----		-----		-----		:
TrCHSa29 :	-----		-----		-----		:
TrCHSa30 :	-----		-----		-----		:
TrCHSa31 :	-----		-----		-----		:
TrCHSa32 :	-----		-----		-----		:
TrCHSa33 :	-----		-----		-----		:
TrCHSa34 :	-----		-----		-----		:
TrCHSa35 :	-----		-----		-----		:
TrCHSa36 :	-----		-----		-----		:
TrCHSa37 :	-----		-----		-----		:
TrCHSa38 :	-----		-----		-----		:
TrCHSa39 :	ATTGTATCAAAGAACATTAATAAAGCATTGGTCGAGGCTTTCCAACCATTAGGAATTTCT						: 541

**FIGURE 15 (cont)**



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	*	1100	*	1120	*	
TrCHSa1	:	-----		-----	:	-
TrCHSa2	:	-----		-----	:	-
TrCHSa3	:	-----		-----	:	-
TrCHSa4	:	-----		-----	:	-
TrCHSa5	:	-----		-----	:	-
TrCHSa6	:	-----		-----	:	-
TrCHSa7	:	-----		-----	:	-
TrCHSa8	:	-----		-----	:	-
TrCHSa9	:	-----		-----	:	-
TrCHSa10	:	-----		-----	:	-
TrCHSa11	:	-----		-----	:	-
TrCHSa12	:	-----		-----	:	-
TrCHSa13	:	-----		-----	:	-
TrCHSa14	:	-----		-----	:	-
TrCHSa15	:	-----		-----	:	-
TrCHSa16	:	-----		-----	:	-
TrCHSa17	:	-----		-----	:	-
TrCHSa18	:	-----		-----	:	-
TrCHSa19	:	-----		-----	:	-
TrCHSa20	:	-----		-----	:	-
TrCHSa21	:	-----		-----	:	-
TrCHSa22	:	-----		-----	:	-
TrCHSa23	:	-----		-----	:	-
TrCHSa24	:	-----		-----	:	-
TrCHSa25	:	-----		-----	:	-
TrCHSa26	:	-----		-----	:	-
TrCHSa27	:	-----		-----	:	-
TrCHSa28	:	-----		-----	:	-
TrCHSa29	:	-----		-----	:	-
TrCHSa30	:	-----		-----	:	-
TrCHSa31	:	-----		-----	:	-
TrCHSa32	:	-----		-----	:	-
TrCHSa33	:	-----		-----	:	-
TrCHSa34	:	-----		-----	:	-
TrCHSa35	:	-----		-----	:	-
TrCHSa36	:	-----		-----	:	-
TrCHSa37	:	-----		-----	:	-
TrCHSa38	:	-----		-----	:	-
TrCHSa39	:	<b>GATTACA</b>	<b>ACTCAATCTTTTGGATTGCACACCCGGGTGGACCTGCAATTCT</b>		:	591

**FIGURE 15 (cont)**

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```

      *           20           *           40           *           60
TrCHSb : TCTTCGNCNAGCTGGACNAACATTTNTGCTTCTTAAAGATGTTTCCTGAGATTGTCTCAAA : 60

      *           80           *           100          *           120
TrCHSb : GAACATTGATAAGGCATTGGTTGAGGCATTCCAACCATTAAACATCTCTGATTACAATTC : 120

      *           140          *           160          *           180
TrCHSb : AATCTTTTGGATTGCTCATCCAGGTGGTCCTGCAATTCTAGACCAAGTTGAGATAAAGTT : 180

      *           200          *           220          *           240
TrCHSb : GGGCTTAAACCTGAAAAAATGAAGGCCACCAGAGATGTACTTAGTGAATATGGTAACAT : 240

      *           260          *           280          *           300
TrCHSb : GTCAAGTGCATGTGTATTGTTTCATCTTAGATGAGATGCAAAGAAATCGGCTGAAAAATGG : 300

      *           320          *           340          *           360
TrCHSb : ACTGAAAACACAGGAGAAGGACTTGACTGGGGTGTGTTGTTTGGATTTGGACCAGGACT : 360

      *           380          *           400          *           420
TrCHSb : TACCATTGAAACTGTTGTTCTACATAGTGTGGCTATATGAGAATGCGAGACTTGATTGTT : 420

      *           440          *           460          *           480
TrCHSb : TTGTATTGTATTGTATTGTATTGTATTACTTTTAATCTTGCTTGAATTTCCATTTAACAA : 480

      *           500          *           520          *           540
TrCHSb : TAAATATGGAGTTCAATAAGTACCATCAGTGTTAAAATAATATATCGTTAATAGCTATTA : 540

      *           560          *           580          *           600
TrCHSb : TTTTAGTGTCTGTTTCTTTTACTAAACTATATTTTATTTTAGTATTTGCTATTGATTTG : 600

      *           620          *
TrCHSb : AAATAAATATTGTCCTCTTAACTGAAAAAAAAA : 634

```

## FIGURE 16

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TrCHSb : LRXAGXTFXLLKDVPEIVSKNIDKALVEAFQPLNISDYNSIFWIAHPGGPAILDQVEIKL : 60

TrCHSb : GLKPEKMKATRDVLSEYGNMSSACVLFILDEMOKKSAENGLKTTGEGLDWGVLFQFGPGGL : 120

TrCHSb : TIETVVLHSVAI : 132

**FIGURE 17**

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```

      *           20           *           40           *           60
TrCHSb1: TCTTCGNCNAGCTGGACNAACATTTNTGCTTCTTAAAGATGTTCCCTGAGATTGTCTCAAA : 60
TrCHSb2: ----- : -
TrCHSb3: ----- : -

      *           80           *           100          *           120
TrCHSb1: GAACATTGATAAGGCATTGGTTGAGGCATTCCAACCATTAACATCTCTGATTACAATTC : 120
TrCHSb2: ----- : -
TrCHSb3: ----- : -

      *           140          *           160          *           180
TrCHSb1: AATCTTTTGGATTGCTCATCCAGGTGGTCCTGCAATTCTAGACCAAGTTGAGATAAAGTT : 180
TrCHSb2: ----- : -
TrCHSb3: ----- : -

      *           200          *           220          *           240
TrCHSb1: GGGCTTAAACCTGAAAAAATGAAGGCCACCAGAGATGTACTTAGTGAATATGGTAACAT : 240
TrCHSb2: ----- : -
TrCHSb3: ----- : -

      *           260          *           280          *           300
TrCHSb1: GTCAAGTGCATGTGTATTGTTTCATCTTAGATGAGATGAGAAAGAAATCGGCTGAAAATGG : 300
TrCHSb2: ----- GAGATGCACAAGAAATCGGCTTAAAATGG : 29
TrCHSb3: ----- GAGATGCCAAGAAATCGGCTCAAAATGG : 29

      *           320          *           340          *           360
TrCHSb1: ACTTAAAACACAGGAGAAGGACTTGACTGGGGTGTGTTGTTTGGATTGTTGGCCCGGACT : 360
TrCHSb2: ACTGAAAACACAGGAGAAGGACTTGACTGGGGTGTGTTGTTTGGATTGTTGGACCAGGACT : 89
TrCHSb3: ACTGAAAACACAGGAGAAGGACTTGACTGGGGTGTGTTGTTTGGATTGTTGGACCAGGACT : 89

      *           380          *           400          *           420
TrCHSb1: TACCATTGAAACTGTTGTTCTACATAGTGTGGCTATATGAGAATGAGACTTGATTGTT : 420
TrCHSb2: TACCATTGAAACTGTTGTTCTACATAGTGTGGCTATATGAGAATGCGAGACTTGATTGTT : 149
TrCHSb3: TACCATTGAAACTGTTGTTCTACATAGTGTGGCTATATGAGAATGCGAGACTTGATTGTT : 149

      *           440          *           460          *           480
TrCHSb1: T--T--T--T--ATTGATTGTATTACTTTTAACTCTGCTTGAAGTTCCATTTAANAA : 470
TrCHSb2: TTGTATTGTATTGTATTGTATTGTATTACTTTTAACTCTGCTTGAAGTTCCATTTAACAA : 209
TrCHSb3: TTGTATTGTATTGTATTGTATTGTATTACTTTTAACTCTGCTTGAAGTTCCATTTAACAA : 209

      *           500          *           520          *           540
TrCHSb1: TAAATATGGNGNTCANTTGGN----- : 491
TrCHSb2: TAAATATGGAGTTCAATAAGTACCATCAGTGTAAATAATATATCGTTAATAGCTATTA : 269
TrCHSb3: TAAATATGGAGTTCAATAAGTACCATCAGTGTAAATAATATATCGTTAATAGCTATTA : 269

      *           560          *           580          *           600
TrCHSb1: ----- : -
TrCHSb2: TTTTAGTGTCTGTTTCTTTTACTAAACTATATTTTATTTTAGTATTGCTATTGATTG : 329
TrCHSb3: TTTTAGTGTCTGTTTCTTTTACTAAACTATATTTTATTTTAGTATTGCTATTGATTG : 329

      *           620          *
TrCHSb1 : ----- : -
TrCHSb2 : AAATAAATATTGTCCTCTTAACTGAAAAAAAAA : 363
TrCHSb3 : AAATAAATATTGTCCTCTTAACTGAAAAAAAAA : 363

```

FIGURE 18

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```

      *           20           *           40           *           60
TrCHSc : GNTTCAATCTGTTGTGCATAAAAATTNCTTTGCNATAGAAAACCATACACATTTGATCTTG : 60

      *           80           *           100          *           120
TrCHSc : CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 120

      *           140          *           160          *           180
TrCHSc : GGGAAGGCTACTATATTGGCTCTTGGCAAGGCATTCCCTCACCAACTTGTGATGCAAGAG : 180

      *           200          *           220          *           240
TrCHSc : TATTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACCTTAAGCAGAAA : 240

      *           260          *           280          *           300
TrCHSc : CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 300

      *           320          *           340          *           360
TrCHSc : ATACTAAAGAAATATCCAGAACTTGTGTGCGAAGGCGCCTCAACTGTAAACAACGTTTA : 360

      *           380          *           400          *           420
TrCHSc : GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG : 420

      *           440          *           460          *           480
TrCHSc : AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT : 480

      *           500          *           520          *           540
TrCHSc : AGATTACCCGGTGGTGACCTATACTTGTCAAAAGGACTAGGACTAAACCCTAAAAATTCAA : 540

      *           560          *           580          *           600
TrCHSc : AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA : 600

      *           620          *           640          *           660
TrCHSc : GACATAGCTGAGAACAACCCTGGAAGTAGAGTTTGTGCTTACTTCTGAAACTACAATT : 660

      *           680          *           700          *           720
TrCHSc : ATTGGATTCAAGCCACCAAGTGTGATAGACCTTATGATCTTGTGTTGGTGTGGCACTCTTT : 720

      *           740          *           760          *           780
TrCHSc : GGAGATGGTGCTGGTGCTATGATAATTGGCTCAGACCCAATACTTGAAACTGAGACTCCA : 780

      *           800          *           820          *           840
TrCHSc : TTGTTTGAGCTTCATACTTCAGCTCAGGAGTTTATACCAGACACAGAGAAGAAAATAGAT : 840

      *           860          *           880          *
TrCHSc : GGGCGGCTGACGGAGGAGGGCATAAGTTTCACGCTAGCGAGGGAAGTCCCGCAGATA : 897

```

## FIGURE 19

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TrCHSc : MGDEGIVRGVTKQTTPGKATILALGKAFPHQLVMQEYLVLDGYFRDTNCDNPELKQKLARL : 60

TrCHSc : CKTTTVKTRYVVMNNEEILKKYPELVVEGASTVKQRLEICNEAVTQMAIEASQVCLKNWGR : 120

TrCHSc : SLSDITHVYVSSSEARLPGGDLVLSKGLGLNPKIQRTMLYFSGCSGGVAGLRVAKDIAE : 180

TrCHSc : NNPGRVLLATSETTIIGFKPPSVDRPYDLVGVALFGDGAGAMIIGSDPILETETPLFEL : 240

TrCHSc : HTSAQEFIPDTEKKIDGRLTEEGISFTLARELPQI : 275

**FIGURE 20**

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```

      *           20           *           40           *           60
TrCHSc1: GNTTCAATCTGTTGTCATAAAATTNCTTTGTCNATAGAAAACNCTACACATTTGATCTTG : 60
TrCHSc2: ---TCAATCTGTTGNGCATTTTNTTNCCTTTGTCNATAGAAAACCATACACATTTGATCTTG : 57
TrCHSc3: -----TCTGTTGTCNTTAACTTNCCTTTGTCNATAGAAAACATACACATTTGATCTTG : 53
TrCHSc4: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : 53
TrCHSc5: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : 40
TrCHSc6: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : 30
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : -

      *           80           *           100          *           120
TrCHSc1: CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 120
TrCHSc2: CTTTAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 117
TrCHSc3: CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 113
TrCHSc4: CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 113
TrCHSc5: CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 100
TrCHSc6: CAAAGAAGAAATATGGGAGACGAAGGTATAGTGAGAGGTGTCACAAAGCAGACAACCCCT : 90
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : -

      *           140          *           160           *           180
TrCHSc1: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 180
TrCHSc2: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 177
TrCHSc3: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 173
TrCHSc4: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 173
TrCHSc5: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 160
TrCHSc6: GGGGAAGGCTACTATATTGGCTCTTTGGCAAGGCATTCCCTCACCACCTTGTGATGCAAGAG : 150
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : -

      *           200          *           220           *           240
TrCHSc1: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 240
TrCHSc2: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 237
TrCHSc3: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 233
TrCHSc4: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 233
TrCHSc5: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 220
TrCHSc6: TTTTAGTTGATGGTTATTTTAGGGACACTAATTGTGACAATCCTGAACTTAAGCAGAAA : 210
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : -

      *           260          *           280           *           300
TrCHSc1: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 300
TrCHSc2: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 297
TrCHSc3: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 293
TrCHSc4: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 293
TrCHSc5: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 280
TrCHSc6: CTTGCTAGACTTTGTAAGACAACCACGGTAAAAACAAGGTATGTTGTTATGAATGAGGAG : 270
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : -

      *           320          *           340           *           360
TrCHSc1: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 360
TrCHSc2: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 357
TrCHSc3: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 353
TrCHSc4: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 353
TrCHSc5: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 340
TrCHSc6: ATACTAAAGAAATATCCAGAACTTGTGTCGAAGGCGCCTCAACTGTAAAAACAACGTTTA : 330
TrCHSc7: -----TCTGTTGTCNTTAAATTACTTTGTCNATAGAAAACNCTACACATTTGATCTTG : 47

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FIGURE 21

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		*	380	*	400	*	420	
TrCHSc1:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 420
TrCHSc2:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 417
TrCHSc3:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 413
TrCHSc4:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 413
TrCHSc5:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 400
TrCHSc6:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 390
TrCHSc7:	GAGATATGTAATGAGGCAGTAACACAAATGGCAATTGAAGCTTCCCAAGTTTGCCTAAAG							: 107
		*	440	*	460	*	480	
TrCHSc1:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 480
TrCHSc2:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 477
TrCHSc3:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 473
TrCHSc4:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 473
TrCHSc5:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 460
TrCHSc6:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 450
TrCHSc7:	AATTGGGGTAGATCCTTATCGGACATAACTCATGTGGTTTATGTTTCATCTAGTGAAGCT							: 167
		*	500	*	520	*	540	
TrCHSc1:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 540
TrCHSc2:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 537
TrCHSc3:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 533
TrCHSc4:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 533
TrCHSc5:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 520
TrCHSc6:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 510
TrCHSc7:	AGATTACCCGGTGGTGACCTATACCTTGTCAAAGGACTAGGACTAAACCCCTAAAAATTCAA							: 227
		*	560	*	580	*	600	
TrCHSc1:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCG-----							: 577
TrCHSc2:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA							: 597
TrCHSc3:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA							: 581
TrCHSc4:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA							: 588
TrCHSc5:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCG-----							: 563
TrCHSc6:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA							: 570
TrCHSc7:	AGAACCATGCTCTATTTCTCTGGATGCTCGGGAGGCGTAGCCGGCCTTCGCGTTGCGAAA							: 287
		*	620	*	640	*	660	
TrCHSc1:	-----							: -
TrCHSc2:	GA-----							: 599
TrCHSc3:	-----							: -
TrCHSc4:	-----							: -
TrCHSc5:	-----							: -
TrCHSc6:	GACATAGCTGAGAACAACCCCTGGAAGTAGAGTT-----							: 603
TrCHSc7:	GACATAGCTGAGAACAACCCCTGGAAGTAGAGTTTGGCTTGCTACTTCTGAAACTACAATT							: 347
		*	680	*	700	*	720	
TrCHSc1:	-----							: -
TrCHSc2:	-----							: -
TrCHSc3:	-----							: -
TrCHSc4:	-----							: -
TrCHSc5:	-----							: -
TrCHSc6:	-----							: -
TrCHSc7:	ATTGGATTCAAGCCACCAAGTGTGATAGACCTTATGATCTTGTGGTGTGGCACTCTTT							: 407

FIGURE 21 (cont)



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	*	740	*	760	*	780	
TrCHSc1:	-----		-----		-----		:
TrCHSc2:	-----		-----		-----		:
TrCHSc3:	-----		-----		-----		:
TrCHSc4:	-----		-----		-----		:
TrCHSc5:	-----		-----		-----		:
TrCHSc6:	-----		-----		-----		:
TrCHSc7:	GGAGATGGTGCTGGTGCTATGATAATTGGCTCAGACCCAATACTTGAAACTGAGACTCCA						: 467

	*	800	*	820	*	840	
TrCHSc1:	-----		-----		-----		:
TrCHSc2:	-----		-----		-----		:
TrCHSc3:	-----		-----		-----		:
TrCHSc4:	-----		-----		-----		:
TrCHSc5:	-----		-----		-----		:
TrCHSc6:	-----		-----		-----		:
TrCHSc7:	TTGTTTGAGCTTCATACTTCAGCTCAGGAGTTTATACCAGACACAGAGAAGAAAATAGAT						: 527

	*	860	*	880	*		
TrCHSc1:	-----		-----		-----	:	-
TrCHSc2:	-----		-----		-----	:	-
TrCHSc3:	-----		-----		-----	:	-
TrCHSc4:	-----		-----		-----	:	-
TrCHSc5:	-----		-----		-----	:	-
TrCHSc6:	-----		-----		-----	:	-
TrCHSc7:	GGGCGGCTGACGGAGGAGGGCATAAGTTTCACGCTAGCGAGGGAAGTCCCGCAGATA					:	584

FIGURE 21 (cont)

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      *           20           *           40           *           60
TrCHSd : GTAGCAACACACACTTTGATTTCTTTTGGAGTCCTTGCTACGTGGCNTTACCAAAAAACG : 60

      *           80           *           100          *           120
TrCHSd : TTGCTAAGTCATCAACCATTCCAATTCTTAATATAACCTATCAGTACTCACCATCTTTT : 120

      *           140          *           160          *           180
TrCHSd : CTCCTCCCTGCTAACTTTATACTTAGAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 180

      *           200          *           220          *           240
TrCHSd : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 240

      *           260          *           280          *           300
TrCHSd : GTGTTGATCAGAGTACATACCCCGACTACTACTTCCGCATCACAAACAGTGAGCACAAGA : 300

      *           320          *           340          *           360
TrCHSd : CAGAGCTCAAAGAAAAATTCCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA : 360

      *           380          *           400          *           420
TrCHSd : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT : 420

      *           440          *           460          *           480
TrCHSd : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG : 480

      *           500          *           520          *           540
TrCHSd : CAACAAAGGCAATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 540

      *           560          *           580          *           600
TrCHSd : GCACCACAAGTGGTGTGGACATGCCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 600

      *           620          *           640          *           660
TrCHSd : TTCGTCCGCATGTGAAGCGTTACATGATGTACCAACAAGGTTGTTTTGCTGGTGGCACGG : 660

      *           680          *           700          *           720
TrCHSd : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAGGTGCCCCGTGTATTGGTGGTTT : 720

      *
TrCHSd : GTTCAGAGATAACTG : 735

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## FIGURE 22

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TrCHSd : MVKVNEIRQAQRAEGPATVLAIGTATPPNCVDQSTYPDYYFRITNSEHKTELKEKFQRM C : 60

TrCHSd : DKSMIKKRYMHLTEEILKENPSLCEYMAPSLDARQDMVVVEVPRLGKEAATKAIKEWGQP : 120

TrCHSd : KSKITHLIFCTTSGVDMPGADYQLTKLLGLRPHVKRYMMYQQGCFAGGTVLRRLAKDLAEN : 180

TrCHSd : NKGARVLVVCSEIT : 194

**FIGURE 23**

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      *           20           *           40           *           60
TrCHSd1 : GTAGCAACACACACTTTGATTTCTTTTGGAGTCCTTGCTACGTGGCNTTACCAAAAAACG : 60
TrCHSd2 : GTAGCAACACACACTTTGATTTCTTTTGGAGTCCTTGCTACGTGGCNTTACCAAAAAACG : 60
TrCHSd3 : GTAGCAACACACACTTTGATTTCTTTTGGAGTCCTTGCTACGTGGCNTTACCAAAAAACG : 60
TrCHSd4 : -----NNNCACNCACACTTTTTTCNATCCCTGCTACGTGGCNTTACCAAAAAACG : 50
TrCHSd5 : ----- : -
TrCHSd6 : ----- : -
TrCHSd7 : ----- : -
TrCHSd8 : ----- : -
TrCHSd9 : ----- : -
TrCHSd10 : ----- : -
TrCHSd11 : ----- : -

      *           80           *           100          *           120
TrCHSd1 : TTGCTTAAGTCATCAACCATTCCAATTCCCTTAATATAACCTATCAGTACTTACCATCTTTT : 120
TrCHSd2 : TTGCTTAAGTCATCAACCATTCCAATTCCCTTAATATAACCTATCAGTACTCACCATCTTTT : 120
TrCHSd3 : TTGCTAAGTCATCAACCATTCCAATTCCCTTAATATAACCTATCAGTACTCACCATCTTTT : 120
TrCHSd4 : TTGCTAAGTCATCAACCATTCCAATTCCCTTAATATAACCTATCAGTACTCACCATCTTTT : 110
TrCHSd5 : -----TTCNATCCCT : 10
TrCHSd6 : -----TTCNATCCCT : 10
TrCHSd7 : -----TTCNAT : 6
TrCHSd8 : ----- : -
TrCHSd9 : ----- : -
TrCHSd10 : ----- : -
TrCHSd11 : ----- : -

      *           140          *           160          *           180
TrCHSd1 : CTTCCCTCCCTGCTAACTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 180
TrCHSd2 : CTTCCCTCCCTGCTAACTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 180
TrCHSd3 : CTTCCCTCCCTGCTAACTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 180
TrCHSd4 : CTTCCCTCCCTGCTAACTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 170
TrCHSd5 : AATTGAGATCTTCTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 70
TrCHSd6 : AATTGAGATCTTCTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 70
TrCHSd7 : AATTGAGATCTTCTTTTAACTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 66
TrCHSd8 : -----CCNNTAACTTTTANAATTNNGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 53
TrCHSd9 : -----TNNNTTTTANAATTNNGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 48
TrCHSd10 : -----ATTCTTCTTGAAGAAGATGGTGAAAGTTAATGAGATCCGCCAGG : 45
TrCHSd11 : ----- : -

      *           200          *           220          *           240
TrCHSd1 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 240
TrCHSd2 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 240
TrCHSd3 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 240
TrCHSd4 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 230
TrCHSd5 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 130
TrCHSd6 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 130
TrCHSd7 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 126
TrCHSd8 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 113
TrCHSd9 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 108
TrCHSd10 : CACAGAGAGCTGAAGGCCCTGCCACTGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 105
TrCHSd11 : -----TAAAACT : 7

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FIGURE 24

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          *           260           *           280           *           300
TrCHSd1 : GTGTCGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :300
TrCHSd2 : GTGTCGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :300
TrCHSd3 : GTGTCGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :300
TrCHSd4 : GTGTTGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :290
TrCHSd5 : GTGTTGATCAGAGTACATACCCCGACTACTATTTC CGAATCACAAACAGCGAACACAAGA :190
TrCHSd6 : GTGTTGATCAGAGTACATACCCCGACTACTATTTC CGAATCACAAACAGCGAACACAAGA :190
TrCHSd7 : GTGTTGATCAGAGTACATACCCCGACTACTATTTC CGAATCACAAACAGCGAACACAAGA :186
TrCHSd8 : GTGTTGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :173
TrCHSd9 : GTGTTGATCAGAGTACATACCCG GACTACTACTTCCGCATCACAAACAGTGAGCACAAGA :168
TrCHSd10: GTGTTGATCAGAGTACATACCCCGACTACTATTTC CGAATCACAAACAGCGAACACAAGA :165
TrCHSd11: GTGTTGATCAGAGTACATACCCCGACTACTATTTC CGAATNNCAAACAGCGAACACAAGA : 67

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          *           320           *           340           *           360
TrCHSd1 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :360
TrCHSd2 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :360
TrCHSd3 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :360
TrCHSd4 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :350
TrCHSd5 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :250
TrCHSd6 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :250
TrCHSd7 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :246
TrCHSd8 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :233
TrCHSd9 : CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :228
TrCHSd10: CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :225
TrCHSd11: CAGAGCTCAAAGAAAAAATTCAGCGCATGTGTGACAAATCTATGATTAAGAAGAGATACA :127

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          *           380           *           400           *           420
TrCHSd1 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :420
TrCHSd2 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :420
TrCHSd3 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :420
TrCHSd4 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :410
TrCHSd5 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :310
TrCHSd6 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :310
TrCHSd7 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :306
TrCHSd8 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :293
TrCHSd9 : TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :288
TrCHSd10: TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :285
TrCHSd11: TGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAAGTTTATGTGAGTACATGGCACCTT :187

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          *           440           *           460           *           480
TrCHSd1 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :480
TrCHSd2 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :480
TrCHSd3 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :480
TrCHSd4 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :470
TrCHSd5 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :370
TrCHSd6 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :370
TrCHSd7 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :366
TrCHSd8 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :353
TrCHSd9 : CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :348
TrCHSd10: CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :345
TrCHSd11: CATTGGATGCAAGACAAGACATGGTGGTTGTGGAAGTACCAAGGCTAGGAAAAGAGGCAG :247

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FIGURE 24 (cont)

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      *           500           *           520           *           540
TrCHSd1 : CAACAAAGGCAATTAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 540
TrCHSd2 : CAACAAAGGCTATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACTCACCTCATCTTTT : 540
TrCHSd3 : CAACAAAGGCTATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACTCACCTCATCTTTT : 540
TrCHSd4 : TAACAAAGGCAATTAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 530
TrCHSd5 : CAACAAAGGCTATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACTCACCTCATCTTTT : 430
TrCHSd6 : CAACAAAGGCTATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACTCACCTCATCTTTT : 430
TrCHSd7 : CAACAAAGGCTATCAAGGAATGGGGTCAACCTAAGTCCAAGATTACTCACCTCATCTTTT : 426
TrCHSd8 : CAACAAAGGCAATTAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 413
TrCHSd9 : CAACAAAGGCAATTAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 408
TrCHSd10 : CAACAAAGGCAATTAAGGAATGGGGTCAACCTAAGTCCAAGATTACCCACCTCATCTTTT : 405
TrCHSd11 : CCN----- : 250

      *           560           *           580           *           600
TrCHSd1 : GCACCACAGTGGTGTGACATGCCCGGTGCCGC----- : 574
TrCHSd2 : GCACCACAAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAG----- : 597
TrCHSd3 : GCACCACAAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 600
TrCHSd4 : GCACCACAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 585
TrCHSd5 : GCACCACAAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 490
TrCHSd6 : GCACCACAAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 490
TrCHSd7 : GCACCACAAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 486
TrCHSd8 : GCACCACAGTGGTGTGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 473
TrCHSd9 : GCACCACAGTGGTGTGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 468
TrCHSd10 : GCACCACAGTGGTGTGGACATGCCCGGTGCCGACTATCAGCTTACAAAGCTTTTAGGCC : 465
TrCHSd11 : ----- : -

      *           620           *           640           *           660
TrCHSd1 : ----- : -
TrCHSd2 : ----- : -
TrCHSd3 : TTCGTCCGCATGTGAN----- : 616
TrCHSd4 : ----- : -
TrCHSd5 : TTCGTCCGCATGTGAAGCGTTATATGATGTACCAACAAGGTTGTTTTGCTGGTGGCACGG : 550
TrCHSd6 : TTCGTCCGCATGTGAAGCGTTATATGATGTACCAACAAGGTTGTTTTGCTGGTGGCACGG : 550
TrCHSd7 : TTCGTCCGCATGTGAAGCGTTATATGATGTACCAACAAGGTTGTTTTGCTGGTGGCACGG : 546
TrCHSd8 : TTCGTCCATATGTGAAGCGTTACATGATGTATCAACAAGGTTGTTTTGCTGGTGGCACGG : 533
TrCHSd9 : TSCGTCCATATGTGAAGCGTTACATGATGTATCAACAAGGTTGTTTTGCTGGTGGCACGG : 528
TrCHSd10 : TSCGTCCATATGTGAAGCGTTACATGATGTATCAACAAGGTTGTTTTGCTGGTGGCACGG : 525
TrCHSd11 : ----- : -

      *           680           *           700           *           720
TrCHSd1 : ----- : -
TrCHSd2 : ----- : -
TrCHSd3 : ----- : -
TrCHSd4 : ----- : -
TrCHSd5 : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAGGTGCCCG----- : 596
TrCHSd6 : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAGGTGCCCGTGTATTTGGTGGTTT : 610
TrCHSd7 : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAGGTGCCCGTGTATTTGGTGGTTT : 606
TrCHSd8 : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAN----- : 571
TrCHSd9 : TGCTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAANGTGCCCGTGTGTTGGTGGTTT : 588
TrCHSd10 : TACTTCGTTTGGCTAAAGACTTGGCTGAAAACAACAAAGGTGCCCGTGTGTTGGTGGTTT : 585
TrCHSd11 : ----- : -

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FIGURE 24 (cont)

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\*

TrCHSd1	:	-----	:	-
TrCHSd2	:	-----	:	-
TrCHSd3	:	-----	:	-
TrCHSd4	:	-----	:	-
TrCHSd5	:	-----	:	-
TrCHSd6	:	<b>G TTCAGAG</b> -----	:	618
TrCHSd7	:	<b>GTT</b> -----	:	609
TrCHSd8	:	-----	:	-
TrCHSd9	:	<b>G TTCANAGATAACTG</b>	:	603
TrCHSd10	:	<b>GTT</b> -----	:	588
TrCHSd11	:	-----	:	-

**FIGURE 24 (cont)**

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TrCHSe : GNAGCAACACACACTTTGATTTCTTTTGAATCCCTGCTACGTGGCNCACCAAAAAACGT : 60  
 TrCHSe : TGCTAAGTCATCAACCATTCCAATTCTTAATATAACCTATCAGTACTCACCATCTTTTC : 120  
 TrCHSe : TTCCTCCCTGCTAACTTTAGACTCAGTAGAAGATGGTGAATGTTAATGAGATCCGCCAGG : 180  
 TrCHSe : CACAGAGAGCTGAAGGCCCTGCCACCGTGTTGGCAATCGGCACTGCAACTCCTCCAAACT : 240  
 TrCHSe : GTGTTGATCAGAGTACATACCCGGACTACTACTTCCGCATCACAAACAGTGAGCACAAGA : 300  
 TrCHSe : CAGAGCTCAAAGAAAAATTCCAGCGCACGTGTAAGATATTTATCTTATACTCCATGCATG : 360  
 TrCHSe : TCTTTTTCTGCTGACTGCCGTGTTTATATATTGTTTTGTTTTGTTTCCTTAAATTTGTTAT : 420  
 TrCHSe : GTCACTCTCACATGTACAAAACACTTAAGACTAACTGCATATCATTTTTTTCAGGGACA : 480  
 TrCHSe : AATCTATGATTAAGAAGAGATACATGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAA : 540  
 TrCHSe : GTTTATGTGAGNACATGGCACCTTCTTGGGATGCAAGACAAGT : 583

## FIGURE 25



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TrCHSe : MVNVNEIRQAQRAEGPATVLAIGTATPPNCVDQSTYPDYYFRITNSEHKTELKEKFQTR : 60

TrCHSe : DKSMIKKRYMHLTEEILKENPSLCEXMAPSWDARQ : 95

**FIGURE 26**

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	*                      20                      *                      40                      *                      60	
TrCHSe1:	GNAGCAACACACACTTTGATTTCTTTTTCGAATCCCTGCTACGTGGCTTACCAAAAAACGT	: 60
TrCHSe2:	-----CTCCCTGCTGCGTGGCNCACC-AAAAACGT	: 29
	*                      80                      *                      100                      *                      120	
TrCHSe1:	TGCTAAGTCATCAACCATTCCAATTCCTTAATATAACCTATCAGTACTCACCATTTTTC	: 120
TrCHSe2:	TGCTAAGT-NTGAACC-TTCC-ATTCCTTAATATAACCTATCAGTACTCACCATCTTTTC	: 86
	*                      140                      *                      160                      *                      180	
TrCHSe1:	TTCTCCCTGCTAACTTTAGACTCAG-AGAAGATGGTGAATGTTAATGAGATCCGCCAGG	: 179
TrCHSe2:	TTCTCCCTGCTAACTTTAGACTCAGTAGAAGATGGTGAATGTTAATGAGATCCGCCAGG	: 146
	*                      200                      *                      220                      *                      240	
TrCHSe1:	CACAGAGAGCTGAAGGCCCTGCCACCGTGTGGCAATCGGCACTGCAACTCCTCCAAACT	: 239
TrCHSe2:	CACAGAGAGCTGAAGGCCCTG-----	: 167
	*                      260                      *                      280                      *                      300	
TrCHSe1:	GTGTTGATCAGAGTACATACCCGGACTACTTCCGCATCACAAACAGTGAGCACAAAGT	: 299
TrCHSe2:	-----	: -
	*                      320                      *                      340                      *                      360	
TrCHSe1:	CAGAGCTCAAAGAAAAATTCCAGCGCACGTGTAAGATATTTATCTTATACTCCATGCATG	: 359
TrCHSe2:	-----	: -
	*                      380                      *                      400                      *                      420	
TrCHSe1:	TCTTTTCTGCTGACTGCCGTGTTTATATATTGTTTGTGTTTGTTCCTTAAATTGTTAT	: 419
TrCHSe2:	-----	: -
	*                      440                      *                      460                      *                      480	
TrCHSe1:	GTCACTCTCACATGTACAAAACACTTAAGACTAAACTGCATATCATTTTTTTCAGGGACA	: 479
TrCHSe2:	-----	: -
	*                      500                      *                      520                      *                      540	
TrCHSe1:	AATCTATGATTAAGAAGAGATACATGCATTTGACAGAAGAGATTTTGAAGGAGAATCCAA	: 539
TrCHSe2:	-----	: -
	*                      560                      *                      580	
TrCHSe1 :	GTTTATGTGAGNACATGGCACCTTCTTGGGATGCAAGACAAGT	: 582
TrCHSe2 :	-----	: -

FIGURE 27

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```

      *           20           *           40           *           60
TrCHSf : GCNTAAGCCTTGATTNNTGTTTGTTCCTAACACAAGAAGTAGTGTTTGCTTGAATCTTA : 60

      *           80           *           100          *           120
TrCHSf : AGAAAAAATGCCTCAAGGTGATTTGAATGGAAGTTCCTCGGTGAATGGAGCACGTGCTAG : 120

      *           140          *           160          *           180
TrCHSf : ACGTGCTCCTACTCAGGGAAAGGCAACGATACTTGCATTAGGAAAGGCTTTCCCGCCCA : 180

      *           200          *           220          *           240
TrCHSf : AGTCCTCCCTCAAGAGTGCTTGGTGGAAGGATTCATTCGCGACACTAAGTGTGACGATAC : 240

      *           260          *           280          *           300
TrCHSf : TTATATTAAGGAGAAATTGGAGCGTCTTTGCAAAAACACAAGTGTAAAAACAAGATACAC : 300

      *           320          *           340          *           360
TrCHSf : AGTAATGTCAAAGGAGATCTTAGACAACATCCAGAGCTAGCCATAGATGGAACACCAAC : 360

      *           380          *           400          *           420
TrCHSf : AATAAGGCAAAAGCTTGAAATAGCAAATCCAGCAGTAGTTGAAATGGCAACAAGAGCAAG : 420

      *   *           440          *           460          *           480
TrCHSf : CAAAGATTGCATCAAAGAATGGGGAAGGTCACCTCAAGATATCACACACATAGTCTATGT : 480

      *           500          *           520          *           540
TrCHSf : TTCCTCGAGCGAAATTTCGTCTACCCGGTGGTGACCTTTATCTTGCAAATGAACTCGGCTT : 540

      *           560          *           580          *           600
TrCHSf : AAACAGCGATGTTAATCGCGTAATGCTCTATTTCTCGGTTGCTACGGCGGTGTCACTGG : 600

      *
TrCHSf : CTTACGTGTCGCC : 613

```

## FIGURE 28

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TrCHSf : MPQGD<sup>\*</sup>LN<sup>20</sup>GSSSVNGARARRAP<sup>\*</sup>TQ<sup>40</sup>GKATILALGKAFPAQVLPQECLVEGFIRD<sup>\*</sup>TKCDDTYI<sup>60</sup> : 60

TrCHSf : KEKLERLCKNTTVKTRYTVMSKEILD<sup>\*</sup>NYPELAIDGTPTIRQ<sup>100</sup>LEIANPAVVEMAT<sup>\*</sup>RASKD<sup>120</sup> : 120

TrCHSf : CIKEWGRSPQDITHIVYVSSSEIRLP<sup>\*</sup>GGDLYLANELGLNSDVNRV<sup>160</sup>MLYFLG<sup>\*</sup>CYGGVTGLR<sup>180</sup> : 180

TrCHSf : VA : 182

## FIGURE 29

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		*	20	*	40	*	60	
TrCHSf1:	GCNTAAGCCTTGATTNTTGTGTTTCCTAACACAAGAAGTGTGTTTGCTTGAATCTTA	:	60					
TrCHSf2:	---TAAGCCTTGATTNTTGTGTTTCCTAACACAAGAAGTGTGTTTGCTTGAATCTTA	:	57					
TrCHSf3:	-----TTGATTGTTGTTTTCCTAACACAAGAAGTGTGTTTGCTTGAATCTTA	:	51					
TrCHSf4:	-----GTTTGTTCCTAACACAAGAAGTGTGTTTGCTTGAATCTTA	:	42					
		*	80	*	100	*	120	
TrCHSf1:	AGAAAAAATGCCTCAAGGTGATTTGAATGGAAGTTCCTCGGTGAATGGAGCACGTGCTAG	:	120					
TrCHSf2:	AGAAAAAATGCCTCAAGGTGATTTGAATGGAAGTTCCTCGGTGAATGGAGCACGTGCTAG	:	117					
TrCHSf3:	AGAAAAAATGCCTCAAGGTGATTTGAATGGAAGTTCCTCGGTGAATGGAGCACGTGCTAG	:	111					
TrCHSf4:	AGAAAAAATGCCTCAAGGTGATTTGAATGGAAGTTCCTCGGTGAATGGAGCACGTGCTAG	:	102					
		*	140	*	160	*	180	
TrCHSf1:	ACGTGCTCCTACTCAGGGAAAGGCAACGATACTTGCATTAGGAAAGGCTTTCCCCGCCCA	:	180					
TrCHSf2:	ACGTGCTCCTACTCAGGGAAAGGCAACGATACTTGCATTAGGAAAGGCTTTCCCCGCCCA	:	177					
TrCHSf3:	ACGTGCTCCTACTCAGGGAAAGGCAACGATACTTGCATTAGGAAAGGCTTTCCCCGCCCA	:	171					
TrCHSf4:	ACGTGCTCCTACTCAGGGAAAGGCAACGATACTTGCATTAGGAAAGGCTTTCCCCGCCCA	:	162					
		*	200	*	220	*	240	
TrCHSf1:	AGTCCTCCCTCAAGAGTGCTTGGTGGAAGGATTCATTGCGGACACTAAGTGTGACGATAC	:	240					
TrCHSf2:	GTCTCTCCCTCAAGAGTGCTTGGTGGAAGGATTCATTGCGGACACTAAGTGTGACGATAC	:	237					
TrCHSf3:	GTCTCTCCCTCAAGAGTGCTTGGTGGAAGGATTCATTGCGGACACTAAGTGTGACGATAC	:	231					
TrCHSf4:	AGTCCTCCCTCAAGAGTGCTTGGTGGAAGGATTCATTGCGGACACTAAGTGTGACGATAC	:	222					
		*	260	*	280	*	300	
TrCHSf1:	TTATATTAAGGAGAAATTGGAGCGTCTTTGCAAAAACACAACCTGTAAAAACAAGATACAC	:	300					
TrCHSf2:	TTATATTAAGGAGAAATTGGAGCGTCTTTGCAAAAACACAACCTGTAAAAACAAGATACAC	:	297					
TrCHSf3:	TTATATTAAGGAGAAATTGGAGCGTCTTTGCAAAAACACAACCTGTAAAAACAAGATACAC	:	291					
TrCHSf4:	TTATATTAAGGAGAAATTGGAGCGTCTTTGCAAAAACACAACCTGTAAAAACAAGATACAC	:	282					
		*	320	*	340	*	360	
TrCHSf1:	AGTAATGTCAAAGGAGATCTTAGACAACCTATCCAGAGCTAGCCATAGATGGAACACCAAC	:	360					
TrCHSf2:	AGTAATGTCAAAGGAGATCTTAGACAACCTATCCAGAGCTAGCCATAGATGGAACACCAAC	:	357					
TrCHSf3:	AGTAATGTCAAAGGAGATCTTAGACAACCTATCCAGAGCTAGCCATAGATGGAACACCAAC	:	351					
TrCHSf4:	AGTAATGTCAAAGGAGATCTTAGACAACCTATCCAGAGCTAGCCATAGATGGAACACCAAC	:	342					
		*	380	*	400	*	420	
TrCHSf1:	AATAAGGCAAAAGCTTGAAATAGCAAATCCAGCAGTAGTTGAAATGGCAACAAGAGCAAG	:	420					
TrCHSf2:	AATAAGGCAAAAGCTTGAAATAGCAAATCCAGCAGTAGTTGAAATGGCAACAAGAGCAAG	:	417					
TrCHSf3:	AATAAGGCAAAAGCTTGAAATAGCAAATCCAGCAGTAGTTGAAATGGCAACAAGAGCAAG	:	411					
TrCHSf4:	AATAAGGCAAAAGCTTGAAATAGCAAATCCAGCAGTAGTTGAAATGGCAACAAGAGCAAG	:	402					
		*	440	*	460	*	480	
TrCHSf1:	CAAAGATTGCATCAAAGAATGGGGAAGGTCACCTCAAGATATCACACACATAGTCTATGT	:	480					
TrCHSf2:	CAAAGATTGCATCAAAGAATGGGGAAGGTCACCTCAAGATATCACACACATAGTCTATGT	:	477					
TrCHSf3:	CAAAGATTGCATCAAAGAATGGGGAAGGTCACCTCAAGATATCACACACATAGTCTATGT	:	471					
TrCHSf4:	CAAAGATTGCATCAAAGAATGGGGAAGGTCACCTCAAGATATCACACACATAGTCTATGT	:	462					

FIGURE 30

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```

      *           500           *           520           *           540
TrCHSf1: TTCCTCGAGCGAAATTCGTCTACCCGGTGGTGACCTTTATCTTGCAAATGAACTCGGCTT : 540
TrCHSf2: TTCCTCGAGCGAAATTCGTCTACCCGGTGGTGACCTTTATCTTGCAAATGAACTCGGCTT : 537
TrCHSf3: TTCCTCGAGCGAAATTCGTCTACCCGGTGGTGACCTTTATCTTGCAAATGAACTCGGCTT : 531
TrCHSf4: TTCCTCGAGCGAAATTCGTCTACCCGGTGGTGACCTTTATCTTGCAAATGAACTCGGCTT : 522

      *           560           *           580           *           600
TrCHSf1: AAACAGCGATGTTAATCGCGTAATGCTCTATTTTCCTCGGTTGCTACGGCGGTGTCAGTGG : 600
TrCHSf2: AAACAGCGATGTTAATCGCGTAATGCTCTATTT----- : 570
TrCHSf3: AAACAGCGATGTTAATCGCGTAATGCTCTATTTTCCTCGGTTGCT----- : 575
TrCHSf4: AAACAGCGATGTTAATCGCGTAATGCTCTATTTTCCTCGGTTGCTACGGCGG----- : 573

      *
TrCHSf1 : CTTACGTGTCGCC : 613
TrCHSf2 : ----- : -
TrCHSf3 : ----- : -
TrCHSf4 : ----- : -

```

## FIGURE 30 (cont)

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TrCHSg : GTATACCAAGGTTGTTTTGCTGGTGGCACGGTACTTCGTTTGGCTAAAGACTTGGCTGAA : 60

TrCHSg : AACAAACAAGGTGCCCCGTGTGTTGGTGGTTTGTTCAGAGATAACTGCAGTTACTTTCCGT : 120

TrCHSg : GGACCCAGTGACACTCACCTTGATAGCCTTGTGGGGCAAGCATTGTTTGGAGATGGTGCA : 180

TrCHSg : GCAGCTGTGATTGTTGGTTCAGACCCTTTGCCAGAAGTTGAGAAGCCTTTGTTGAATTG : 240

TrCHSg : GTATGGACCGCACAAACAATCGCTCCAGATAGTGAAGGAGCCATTGATGGTCACCTTCGC : 300

TrCHSg : GAAGCAGGGCTGACATTCCATCTCCTCAAGGATGTTCTTAGCCTTGTCTCAAATAACATT : 360

TrCHSg : GAGAAAGCGCTTGTTGATGCCTTTCAACCTTTGAATATTTCTGACTACAATTCCATCTTT : 420

TrCHSg : TGGATTGCACACCCAGGCGGACCAGCAATTCTTGACCAAGTTGAAGCTAAGTTAGGCTTA : 480

TrCHSg : AAGCCAGAGAAAATGCAAGCCACTCGACATGTACTTAGCGAATATGGTAACATGTCAAGT : 540

TrCHSg : GCGTGTGTGTTATTTATCTTGATGAGATGAGGAGGAAGTCAAAAGAAGACGGACTTGCC : 600

TrCHSg : ACAACAG : 607

## FIGURE 31

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TrCHSg : VYQGC\*FAGGTVLRLAKDLAENNKGARVLVCSEITAVTFRG\*PSDTHLDSL\*VGQALFGDGA : 60

TrCHSg : AAVIVGSDPLPEVEKPLFELVWTAQTIAPDSEGAIDGHLREAGLTFHLLKDVPSLVSNNI : 120

TrCHSg : EKALVDAFQPLNISDYN\*SI\*FWIAHPGGPAILDQVEAKLGLKPEKMQATR\*HVLSEYGNMSS : 180

TrCHSg : ACVLFI\*LD\*EMRRKSKEDGLATT : 202

**FIGURE 32**



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	*                      20                      *                      40                      *                      60	
TrCHSg1:	GTATACCAAGGTTGTTTTGCTGGTGGCACGGTACTTCGTTTGGCTTAAGACTTGGCTGAA	: 60
TrCHSg2:	-----CCAAGGTTGTTTTGCTGGTGGCACGGTACTTCGTTTGGCTAAAGACTTGGCTGAA	: 55
TrCHSg3:	-----GGTTGTTTTGCTGGTGGCACGGTACTTCGTTTGGCTAAAGACTTGGCTGAA	: 51
	*                      80                      *                      100                      *                      120	
TrCHSg1:	AACAACAAAGGTGCCCCGTGTGTTGGTGGTTTGTTCAGAGATAACTGCAGTTACTTTCCGT	: 120
TrCHSg2:	AACAACAAAGGTGCCCCGTGTGTTGGTGGTTTGTTCAGAGATAACTGCAGTTACTTTCCGT	: 115
TrCHSg3:	AACNACAAAGGTGCCCCGTGTGTTGGTGGTTTGTTCAGAGATAACTGCAGTTACTTTCCGT	: 111
	*                      140                      *                      160                      *                      180	
TrCHSg1:	GGACCCAGTGACACTCACTTGTATAGCCTTGTGGGGCAAGCATTTGTTTGGAGATGGTGCA	: 180
TrCHSg2:	GGACCCAGTGACACTCACCTTGTATAGCCTTGTGGGGCAAGCATTTGTTTGGAGATGGTGCA	: 175
TrCHSg3:	GGACCCAGTGACACTCACCTTGTATAGCCTTGTGGGGCAAGCATTTGTTTGGAGATGGTGCA	: 171
	*                      200                      *                      220                      *                      240	
TrCHSg1:	GCAGCTGTGATTGTTGGTTTCAGACCCCTTTGCCAGAAGTTGAGAAGCCTTTGTTTGAATTG	: 240
TrCHSg2:	GCAGCTGTGATTGTTGGTTTCAGACCCCTTTGCCAGAAGTTGAGAAGCCTTTGTTTGAATTG	: 235
TrCHSg3:	GCAGCTGTGATTGTTGGTTTCAGACCCCTTTGCCAGAAGTTGAGAAGCCTTTGTTTGAATTG	: 231
	*                      260                      *                      280                      *                      300	
TrCHSg1:	GTATGGACCGCACAAACAATCGCTCCAGATAGTGAAGGAGCCATTGATGGTCACCTTCGC	: 300
TrCHSg2:	GTATGGACCGCACAAACAATCGCTCCAGATAGTGAAGGAGCCATTGATGGTCACCTTCGC	: 295
TrCHSg3:	GTATGGACCGCACAAACAATCGCTCCAGATAGTGAAGGAGCCATTGATGGTCACCTTCGC	: 291
	*                      320                      *                      340                      *                      360	
TrCHSg1:	GAAGCAGGGCTGACATTCCATCTCCTCAAGGATGTTCCCTAGCCTTGTCTCAAATAACATT	: 360
TrCHSg2:	GAAGCAGGGCTGACATTCCATCTCCTCAAGGATGTTCCCTAGCCTTGTCTCAAATAACATT	: 355
TrCHSg3:	GAAGCAGGGCTGACATTCCATCTCCTCAAGGATGTTCCCTAGCCTTGTCTCAAATAACATT	: 351
	*                      380                      *                      400                      *                      420	
TrCHSg1:	GAGAAAGCGCTTGTGATGCCTTTCAACCTTTGAATATTTCTGACTACAATTCCATCTTT	: 420
TrCHSg2:	GAGAAAGCGCTTGTGATGCCTTTCAACCTTTGAATATTTCTGACTACAATTCCATCTTT	: 415
TrCHSg3:	GAGAAAGCGCTTGTGATGCCTTTCAACCTTTGAATATTTCTGACTACAATTCCATCTTT	: 411
	*                      440                      *                      460                      *                      480	
TrCHSg1:	TGGATTGCACACCCAGGCGGACCAGCAATTCTTGACCAAGTTGAAGCTAAGTTAGGCTTA	: 480
TrCHSg2:	TGGATTGCACACCCAGGCGGACCAGCAATTCTTGACCAAGTTGAAGCTAAGTTAGGCTTA	: 475
TrCHSg3:	TGGATTGCACACCCAGGCGGACCAGCAATTCTTGACCAAGTTGAAGCTAAGTTAGGCTTA	: 471
	*                      500                      *                      520                      *                      540	
TrCHSg1:	AAGCCAGAGAAAATGCAATCCACTCGACATGTACTTAGCGAATATGGTAACATGTCAAGT	: 540
TrCHSg2:	AAGCCAGAGAAAATGCAAGCCACTCGACATGTACTTAGCGAATATGGTAACATGTCAAGT	: 535
TrCHSg3:	AAGCCAGAGAAAATGCAAGCCACTCGACATGTACTTAGCGAATATGGTAACATGTCAAGT	: 531

FIGURE 33

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```

          *           560           *           580           *           600
TrCHSg1: GCGTGTGTGTTATTTATCTTGGATGAGATGAGGAGGAAGTCAAAAGAAGACGGACTTGCC : 600
TrCHSg2: GCGTGTGTGTTATTTATCTTGGATGAGATGAGGAGGAAGTCAAAAGAAGACGGACTTGCC : 591
TrCHSg3: GCGTGTGTGTTATTTATCTTGGATGAGATGAGGAGGAAGTCAAAAGAAGACGGACTTGCC : 590

TrCHSg1 : ACAACAG : 607
TrCHSg2 : ----- : -
TrCHSg3 : ----- : -
```

## FIGURE 33 (cont)

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```

                *           20           *           40           *           60
TrCHSh : AATNACACCNTNANACCTTCCAATTCTCGTACCTCACCAATCTCATTTTTATTATATATC : 60

                *           80           *           100          *           120
TrCHSh : TTGGTACATCTTTTGTTACCTCCAACAAAAAATGGTGACCGTAGAAGAGATTCGTAACG : 120

                *           140          *           160          *           180
TrCHSh : CCCAACGTTCAAATGGCCCTGCCACTATCTTAGCTTTTGGCACAGCCACTCCTTCTAACT : 180

                *           200          *           220          *           240
TrCHSh : GTGTCACCTCAAGCTGATTATCCTGATTACTACTTTCGTATCACCAACAGCGAACATATGA : 240

                *           260          *           280          *           300
TrCHSh : CTGATCTTAAGGAAAAATTCAAGCGGATGTGTGATAGATCAATGATAAAGAAACGTTACA : 300

                *           320          *           340          *           360
TrCHSh : TGCACCTAACAGAAGACTTTCTGAAGGAGAATCCAAATATGTGTGAATACATGGCACCAT : 360

                *           380          *           400          *           420
TrCHSh : CACTAGATGTAAGACGAGACATAGTGGTTGTTGAAGNACCAAAGCTAGGTAAAGAANCAC : 420

                *           440          *           460          *           480
TrCHSh : CAAAAAAGCCATATGNGAATGGGGACAACCAAAATCNAAAATCACACATGCTTGGGTTTC : 480

                *           500          *
TrCHSh : TGACCACTTCCGGTGNTGACATGCCCCGGGG : 510

```

## FIGURE 34

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TrCHSh : MVTVEEIRNAQRSNGPATILAFGTATPSNCVTQADYPDYYFRITNSEHMTDLKEKFKRC : 60

TrCHSh : DRSMIKKRYMHLTEDFLKENPNMCEYMAPSLDVRRDIVVVEPKLGKEPKKAIXEWGQPK : 120

TrCHSh : XKITHAWFLTTSGDMPG : 137

**FIGURE 35**

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TrCHRa : GACAAATGCNTGTGGTTGGAATGGGATCCGCACCTGATTTTACATGTAAGAAAGACACAA : 60

TrCHRa : AAGATGCAATCGTTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGCTT : 120

TrCHRa : ATGGCTCANAACAAGCTCTTGGTGAAGGTTTGAAAGAAGCANTTGAACCTTGGTCTTGTCA : 180

TrCHRa : CTANAGAAGAGCTNTTTGTTACTTCTAACTTTGGGNNACTGAAAATCATNCTAACCTTG : 240

**FIGURE 36**

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TrCHRa : QMXVVGMSAPDFTCKKDTKDAIVEAIKQGYRHFDTAAAYGSXQALGEGLKEAXELGLVT : 60

TrCHRa : XEEXFVTSKLWXTENHXNL : 79

**FIGURE 37**

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TrCHRB : GTGTAGCAGAGTNAGAAAAAGAGAGAAAAAAAACATGGCAGGAAAGAAAATCCCAGAAG : 60

TrCHRB : TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC : 120

TrCHRB : GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC : 180

TrCHRB : ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAG : 240

TrCHRB : CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA : 300

TrCHRB : ATACTGATGCAGATTATGAACTTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG : 360

TrCHRB : GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG : 420

TrCHRB : AAAACCCTGTTGTTTTTCACCAAAGAAGATTACTTCCCTTTGATATAGAAGGGACATGGA : 480

TrCHRB : AAGCTATGGAAGAATGTTATAAGTTAGGCTTAGCAAAGTCTATTGGTATATGCAATTATG : 540

TrCHRB : GTACCAAAAACTCACCAAACTCTTGAAACAGCCACCATTACCCCTGCAGTCAATCAGG : 600

TrCHRB : TGGA : 604

## FIGURE 38

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TrCHRB :                   \*                  20                  \*                  40                  \*                  60  
MAGKKIPEVLLNSGHKMPVIGMGTSVDNRPSNDVLASIFVDAIEVGYRHFDSASVYGTEE :   60

                          \*                  80                  \*                  100                  \*                  120  
TrCHRB : AIGIALAKALEKGLIKSRDEVFITSKPWNTPDADYELIVPALKTTLKKGTEYVDLYLIHW : 120

                          \*                  140                  \*                  160                  \*                  180  
TrCHRB : PVRLRHDLENPVVFTKEDLLPFDIEGTWKAMEECYKLGLAKSIGICNYGTTKLTCLLETA : 180

TrCHRB : TITPAVNQV : 189

**FIGURE 39**



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TrCHRB1:	GTGTAGCAGNATTAGANAAANATAAAAAAAAAAATGGCAGGAAAGAAAAATCCCAGAAG	: 60
TrCHRB2:	--GTAGCAGNGTTAGNANAAGNGAAAAAAAAAATGGCAGGAAAGAAAAATCCCAGAAG	: 58
TrCHRB3:	-----AGTNNGAAAAAGAGAGAAAAAAAAAACNTGGCAGGAAAGAAAAATCCCAGAAG	: 51
TrCHRB4:	-----AGTNNGAAAAAGAGAGAAAAAAAAAACNTGGCAGGAAAGAAAAATCCCAGAAG	: 51
TrCHRB5:	-----ACATGGCAGGAAAGAAAAATCCCAGAAG	: 27
TrCHRB1:	TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC	: 120
TrCHRB2:	TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC	: 118
TrCHRB3:	TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC	: 111
TrCHRB4:	TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC	: 111
TrCHRB5:	TGTTATTGAATTCAGGACACAAAATGCCAGTCATAGGAATGGGAACATCAGTAGACAATC	: 87
TrCHRB1:	GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC	: 180
TrCHRB2:	GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC	: 178
TrCHRB3:	GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC	: 171
TrCHRB4:	GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC	: 171
TrCHRB5:	GTCCATCAAATGATGTTCTTGCTTCAATCTTTGTTGATGCAATTGAAGTTGGTTATCGCC	: 147
TrCHRB1:	ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAAG	: 240
TrCHRB2:	ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAAG	: 238
TrCHRB3:	ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAAG	: 231
TrCHRB4:	ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAAG	: 231
TrCHRB5:	ATTTTCGATTCTGCTTCTGTGTATGGAACAGAGGAAGCCATAGGAATTGCTTTAGCAAAAAG	: 207
TrCHRB1:	CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA	: 300
TrCHRB2:	CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA	: 298
TrCHRB3:	CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA	: 291
TrCHRB4:	CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA	: 291
TrCHRB5:	CTTTAGAAAAAGGGCTTATTAAGAGTAGAGATGAAGTTTTCATCACTTCAAAGCCATGGA	: 267
TrCHRB1:	ATACTGATGCAGATTATGAACCTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG	: 334
TrCHRB2:	ATACTGATGCAGATTATGAACCTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG	: 358
TrCHRB3:	ATACTGATGCAGATTATGAACCTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG	: 351
TrCHRB4:	ATACTGATGCAGATTATGAACCTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG	: 351
TrCHRB5:	ATACTGATGCAGATTATGAACCTATTGTTCCAGCTCTCAAGACCACATTGAAAAAGCTGG	: 327
TrCHRB1:	GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG	: 418
TrCHRB2:	GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG	: 411
TrCHRB3:	GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG	: 411
TrCHRB4:	GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG	: 411
TrCHRB5:	GGACGGAGTATGTGGATCTTTATCTGATCCATTGGCCAGTGAGACTTAGACATGATCTTG	: 334

FIGURE 40

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	*	440	*	460	*	480	
TrCHRB1:	-----						:
TrCHRB2:	AAAACCCTGTTGTTTTCACCAAAGAAGATTTACTTCCCTTTGATATAGAAGGGACATGGA						: 478
TrCHRB3:	AAAACCCTGTTGTTTTCACCAAAGAAGATTTACTTCCCTTTGATATAGAAGGGACATGGA						: 471
TrCHRB4:	AAAACCCTGTTGTTTTCACCAAAGAAGATTTACTTCCCTTTGATATAGAAGGGACATGGA						: 471
TrCHRB5:	-----						:

	*	500	*	520	*	540	
TrCHRB1:	-----						:
TrCHRB2:	AAGCTATGGAAGAATGTTATAAGTTAGGCTTAGCAAAGTCTATTGGTATATGCAATTATG						: 538
TrCHRB3:	AAGCTATGGAAGAATGTTATAAGTTAGGCTTAGCAAAGTCTATTGGTATATGCAATTATG						: 531
TrCHRB4:	AAGCTATGGAAGAATGTTATAAGTTAGGCTTAGCAAAGTCTATTGGTATATGCAATTATG						: 531
TrCHRB5:	-----						:

	*	560	*	580	*	600	
TrCHRB1:	-----						:
TrCHRB2:	GTACCAAAAAAATCACCAAACCTCTTGGAACAGCCACCATTACCCCTGCAGTCAATCAGG						: 598
TrCHRB3:	GTACCAAAAAAATCACCAAACCTCTTGGAACAGCCACCATTACCCCTGCAGTC-----						: 584
TrCHRB4:	GTACCAAAAAAATCAC-----						: 547
TrCHRB5:	-----						:

TrCHRB1:	----	:	-
TrCHRB2:	TGGA	:	602
TrCHRB3:	----	:	-
TrCHRB4:	----	:	-
TrCHRB5:	----	:	-

FIGURE 40 (cont)

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TrCHRC : TAAGAATGAANCAATTTTATCTNANAAAAGGNNCANGCAAGTNAGTTNNATTCAAACATA : 60
          *      20      *      40      *      60

TrCHRC : GNCTTAAAGTGTGTAACATATTCTTAACTTAAANNNTTTTNACCCNACAAAAAAAACAA : 120
          *      80      *      100     *      120

TrCHRC : AGACAATAACATGGGTAGTGTTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 180
          *      140     *      160     *      180

TrCHRC : AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTCACATGTAAGAAAGATAC : 240
          *      200     *      220     *      240

TrCHRC : AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC : 300
          *      260     *      280     *      300

TrCHRC : TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAAGAAGCAATTGAACTTGGCCTTGT : 360
          *      320     *      340     *      360

TrCHRC : CACTAGAGAAGAGCTTTTTTGTACTTCTAAACTTTGGGTCACTGAAAATCATCTCACCT : 420
          *      380     *      400     *      420

TrCHRC : TGTTGTTCTCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA : 480
          *      440     *      460     *      480

TrCHRC : TTTGATCCATTGGCCACTTAGTTCTCAGCCTGGAAAAGTTTTCATTTCCAATTGATGTGGC : 540
          *      500     *      520     *      540

TrCHRC : AGATCTCTTGCCATTTGATGTGAAGGTTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 600
          *      560     *      580     *      600

TrCHRC : TGGACTCACTAAAGCTATTGGTGTAGTAACTTCTCTGTCAAGAACTTCAAATCTTGT : 660
          *      620     *      640     *      660

TrCHRC : CTCAGTTGCCACCGTTCTTCCTGCGGTCAATCAG : 694
          *      680     *

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## FIGURE 41

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TrCHRC : MGSVEIPTKVL<sup>\*</sup>TNTSSQVKMPVVGMGSAPDFTCKKDTKDAIIEAIKQGYRHFD<sup>\*</sup>TAAAYGS : 60

TrCHRC : EQALGEGLKEAIELGLV<sup>\*</sup>TREELFVT<sup>\*</sup>SKLWVTENH<sup>\*</sup>PHLVVPALQKSLKTLQLE<sup>\*</sup>YLDLYLIH : 120

TrCHRC : WPLSSQPGKF<sup>\*</sup>SFPIDVADLLPFDVKG<sup>\*</sup>VWESMEEG<sup>\*</sup>KLGLTKAIGVSNFSV<sup>\*</sup>KKLQNLVSVA : 180

TrCHRC : TVLPAVNQ : 188

**FIGURE 42**

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```

      *           20           *           40           *           60
TrCHRC1 : TAAGAATGAANCAATTTTATCTTATAAAAAGGTCNCANGCAAGTTAGTTTNCATTAAACATA : 60
TrCHRC2 : -----TCTNNNAAAAGGNNCANGCAAGTNAGTTTNTTANTTAACTA : 41
TrCHRC3 : -----GNAATTCAAACATA : 13
TrCHRC4 : -----ATTCAAACATA : 11
TrCHRC5 : -----TTCAAACATA : 10
TrCHRC6 : -----TTCNAAACATA : 10
TrCHRC7 : -----AACATA : 6
TrCHRC8 : -----ATA : 3
TrCHRC9 : -----CA : 2
TrCHRC10 : -----CA : 2
TrCHRC11 : -----CA : 2
TrCHRC12 : -----CA : 2
TrCHRC13 : ----- : -
TrCHRC14 : ----- : -
TrCHRC15 : ----- : -
TrCHRC16 : ----- : -

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      *           80           *           100           *           120
TrCHRC1 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 120
TrCHRC2 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 101
TrCHRC3 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 73
TrCHRC4 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 71
TrCHRC5 : GCCTCCAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 70
TrCHRC6 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 70
TrCHRC7 : GNCTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 66
TrCHRC8 : NTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 63
TrCHRC9 : NTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 62
TrCHRC10 : NTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 62
TrCHRC11 : NTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 62
TrCHRC12 : NTTAAAGTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 62
TrCHRC13 : -----CAGTGTAAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 52
TrCHRC14 : -----CAGTGTAAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 52
TrCHRC15 : -----GTGTGTAACATATTCCTTAACCTAAACNTTTTACCCACAAAAAAAACAA : 52
TrCHRC16 : -----NTTCTTACCTAAACNTTTTACCCACAAAAAAAACAA : 38

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      *           140           *           160           *           180
TrCHRC1 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 180
TrCHRC2 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 161
TrCHRC3 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 133
TrCHRC4 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 131
TrCHRC5 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 130
TrCHRC6 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 130
TrCHRC7 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 126
TrCHRC8 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 123
TrCHRC9 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 122
TrCHRC10 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 122
TrCHRC11 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 122
TrCHRC12 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 122
TrCHRC13 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 112
TrCHRC14 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 112
TrCHRC15 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 112
TrCHRC16 : AGACAATAACATGGGTAGTGTGAAATTCCAACAAAGGTTCTTACTAACACTTCTAGTCA : 98

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FIGURE 43

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	*	200	*	220	*	240	
TrCHRC1 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 240
TrCHRC2 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 221
TrCHRC3 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 193
TrCHRC4 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 191
TrCHRC5 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 190
TrCHRC6 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 190
TrCHRC7 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 186
TrCHRC8 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 183
TrCHRC9 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 182
TrCHRC10 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 182
TrCHRC11 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 182
TrCHRC12 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 182
TrCHRC13 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 172
TrCHRC14 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 172
TrCHRC15 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 172
TrCHRC16 :	AGTGAAAATGCCTGTGGTTGGAATGGGATCAGCTCCTGATTTACATGTAAGAAAAGATAC						: 158

	*	260	*	280	*	300	
TrCHRC1 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 300
TrCHRC2 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 281
TrCHRC3 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 253
TrCHRC4 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 251
TrCHRC5 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 250
TrCHRC6 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 250
TrCHRC7 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 246
TrCHRC8 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 243
TrCHRC9 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 242
TrCHRC10 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 242
TrCHRC11 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 242
TrCHRC12 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 242
TrCHRC13 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 232
TrCHRC14 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 232
TrCHRC15 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 232
TrCHRC16 :	AAAAGATGCAATCATTGAAGCCATCAAACAAGGTTATAGACACTTTGATACTGCTGCTGC						: 218

	*	320	*	340	*	360	
TrCHRC1 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 360
TrCHRC2 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 341
TrCHRC3 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 313
TrCHRC4 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 311
TrCHRC5 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 293
TrCHRC6 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 310
TrCHRC7 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 306
TrCHRC8 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 303
TrCHRC9 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 302
TrCHRC10 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 302
TrCHRC11 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 302
TrCHRC12 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 302
TrCHRC13 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 292
TrCHRC14 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 292
TrCHRC15 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 292
TrCHRC16 :	TTATGGCTCAGAACAAGCTCTTGGTGAAGGTTTGAAGAAGCAATTGAACCTGGCTTGT						: 278

FIGURE 43 (cont)

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		*	380	*	400	*	420	
TrCHRC1 :	CACTAGAGAAGAGCTTTTTGTTGCTTCTAAACTTTGGGTCACTGAAAATCATCCTCACT						: 420	
TrCHRC2 :	CACTAGAGAAGAGCTTTTTGTTGCTTCTAAACTTTGGGTCACTGAAAATCATCCTCACT						: 401	
TrCHRC3 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 373	
TrCHRC4 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 371	
TrCHRC5 :	-----						: -	
TrCHRC6 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 370	
TrCHRC7 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 366	
TrCHRC8 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACT						: 363	
TrCHRC9 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 362	
TrCHRC10 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 362	
TrCHRC11 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACT						: 362	
TrCHRC12 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 362	
TrCHRC13 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 352	
TrCHRC14 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 352	
TrCHRC15 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 352	
TrCHRC16 :	CACTAGAGAAGAGCTTTTTGTTACTTCTAAACTTTGGGTCACTGAAAATCATCCTCACCT						: 338	
		*	440	*	460	*	480	
TrCHRC1 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 480	
TrCHRC2 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 461	
TrCHRC3 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 433	
TrCHRC4 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 431	
TrCHRC5 :	-----						: -	
TrCHRC6 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 430	
TrCHRC7 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 426	
TrCHRC8 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 423	
TrCHRC9 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 422	
TrCHRC10 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 422	
TrCHRC11 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 422	
TrCHRC12 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 422	
TrCHRC13 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 412	
TrCHRC14 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 412	
TrCHRC15 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 412	
TrCHRC16 :	TGTTGTTCCCTGCTCTTCAAAAATCTCTCAAGACTCTTCAATTGGAGTACTTGGACTTGTA						: 398	
		*	500	*	520	*	540	
TrCHRC1 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 540	
TrCHRC2 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 521	
TrCHRC3 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 493	
TrCHRC4 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 491	
TrCHRC5 :	-----						: -	
TrCHRC6 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 490	
TrCHRC7 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 486	
TrCHRC8 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 483	
TrCHRC9 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 482	
TrCHRC10 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 482	
TrCHRC11 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 482	
TrCHRC12 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 482	
TrCHRC13 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 472	
TrCHRC14 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 472	
TrCHRC15 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 472	
TrCHRC16 :	TTTGATCCATTGGCCACTTAGTTCTCAGCCGGAAAGTTTTTCATTTCCAATTGAGGTGGC						: 458	

FIGURE 43 (cont)

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          *           560           *           580           *           600
TrCHRC1 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGG----- : 576
TrCHRC2 : AGN----- : 524
TrCHRC3 : AGATCTCTTGCCATTTGATGTGAGGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 553
TrCHRC4 : AGATCTCTTGCCATTTGATGTGAGGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 551
TrCHRC5 : ----- : -
TrCHRC6 : AGATCTCTTGCCATTTGATGTGAANGGTGTGTTGGGAATCCATGGAAGAAAGCCTTNAAACT : 550
TrCHRC7 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 546
TrCHRC8 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 543
TrCHRC9 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 542
TrCHRC10 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 542
TrCHRC11 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 542
TrCHRC12 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 542
TrCHRC13 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAAGGCTTGAAACT : 532
TrCHRC14 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 532
TrCHRC15 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 532
TrCHRC16 : AGATCTCTTGCCATTTGATGTGAAGGGTGTGTTGGGAATCCATGGAAGAAGGCTTGAAACT : 518

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          *           620           *           640           *           660
TrCHRC1 : ----- : -
TrCHRC2 : ----- : -
TrCHRC3 : TGGACTCACTAAAGCTATTGGTGT----- : 577
TrCHRC4 : TGGACTCACTAAAGCTATTGGTGTAGTAACCTTCTCTGTCAAGAA----- : 597
TrCHRC5 : ----- : -
TrCHRC6 : TGGACTCACTAAAGCTATTGGTGTGCAANNAACCTTCTCTNTCAANAAACTTCAAAATCTTNT : 610
TrCHRC7 : TGGACTCACTAAAGCTATTGGTGT----- : 570
TrCHRC8 : TGGACTCACTAAAGCTATTGGTGTTAGTAACCTTCTCTGTCAAGAAACTTCAAAATCTTGT : 603
TrCHRC9 : TGGACTCACTAAAGCTATTGGTGTTAGTAAN----- : 573
TrCHRC10 : TGGACTCACTAAAGCTATTGGTGTTAGTAACCTTCTCTGTCT----- : 583
TrCHRC11 : TGGACTCACTAAAGCTATTGGTGTTAGTAACCTTCTCTGTCAAGAAACTTCAAAAT----- : 597
TrCHRC12 : TGGACTCACTAAAGCTATTGGTGTTAGTAACCTTCTCTGTCAAGAAACTTCAAAAT----- : 597
TrCHRC13 : TGGACTCACTAAAGCTATTGGTGTTAGTAACCTTCTCTGTCAAGAAACTTCAAAAT----- : 574
TrCHRC14 : TGGACTCACTAAAGCTATTGGTGTAGTAACCTTCTCTGTCAAGAAAN----- : 578
TrCHRC15 : TGGACTCACTAAAGCTATTGGTGTAGTAACCTTCTCTGTCAAGAAACTTCAAAATCTTGT : 592
TrCHRC16 : TGGACTCACTAAAGCTATTGGTGTAGTAACCTTCTCTGTCAAGAAACTTCAAAATCTTGT : 578

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          *           680           *
TrCHRC1 : ----- : -
TrCHRC2 : ----- : -
TrCHRC3 : ----- : -
TrCHRC4 : ----- : -
TrCHRC5 : ----- : -
TrCHRC6 : CTCATTGCCCACCGTTCTTCTGCGG----- : 636
TrCHRC7 : ----- : -
TrCHRC8 : CTCAGTTGCCACCGTTCT----- : 621
TrCHRC9 : ----- : -
TrCHRC10 : ----- : -
TrCHRC11 : ----- : -
TrCHRC12 : ----- : -
TrCHRC13 : ----- : -
TrCHRC14 : ----- : -
TrCHRC15 : CTCAGTTG----- : 600
TrCHRC16 : CTCAGTTGCCACCGTTCTTCTGCGGTCAATCAG : 612

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FIGURE 43 (cont)



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      *           20           *           40           *           60
TrDFRa : GCACACNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 60

      *           80           *           100          *           120
TrDFRa : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTTCGCCTCCTCCTCGAACG : 120

      *           140          *           160           *           180
TrDFRa : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 180

      *           200          *           220           *           240
TrDFRa : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTTCGAAATGGATCTTCTTAACAG : 240

      *           260          *           280           *           300
TrDFRa : CGACTCTATTGCGGCCGCGGTGAAAGGTTGTGCCGGAGTTATACATCTTGCATGTCCTAA : 300

      *           320          *           340           *           360
TrDFRa : CATCATTTGGTGAAGTCAAAGACCCCCGAGAAGCAAATTTTGGAACCGCAATTCAAGGAAC : 360

      *           380          *           400           *           420
TrDFRa : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 420

      *           440          *           460           *           480
TrDFRa : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGAGAAGATTG : 480

      *           500          *           520           *           540
TrDFRa : TTGGACAGACCTTGATTATTGCAAGGAAAAGAAATTATACTACCCCATTTGCAAAGACACT : 540

      *           560          *           580           *           600
TrDFRa : AGCAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGACCGGTTTTTGATGTTGTTATGATTAA : 600

      *
TrDFRa : CCCTGGTACT : 610

```

## FIGURE 44

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TrDFRa : MSKTVCTXGASGAIGSWVVRLLLLERGYIVHATIQDLEDENETKHLEAMEGAKGHLKFFEM : 60

TrDFRa : DLLNSDSIAAAVKGCAGVIHLACPNIIGEVKDPEKQILEPAIQGTNVNLKVAKEAGVERV : 120

TrDFRa : VATSSISAIIPSPNWPADKIKGEDCWTDLDYCKEKKLYYPIAKTLAEKAGWEFAKETGFD : 180

TrDFRa : VVMINPGT : 188

## FIGURE 45

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```

      *           20           *           40           *           60
TrDFRa1 : GCACACCTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 60
TrDFRa2 : GCNCACNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 60
TrDFRa3 : GCACACNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 60
TrDFRa4 : GCACACNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 60
TrDFRa5 : --GCCNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 58
TrDFRa6 : --GCTNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 58
TrDFRa7 : -----CNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 55
TrDFRa8 : -----CNTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 55
TrDFRa9 : -----TTTCTTGACTTACCAATTGTAATNCATAATAATTNTAAACATGTCAAAGACAGT : 54
TrDFRa10 : -----NNGGT : 5

      *           80           *           100           *           120
TrDFRa1 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 120
TrDFRa2 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 120
TrDFRa3 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 120
TrDFRa4 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 120
TrDFRa5 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 118
TrDFRa6 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 118
TrDFRa7 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 115
TrDFRa8 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 115
TrDFRa9 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 114
TrDFRa10 : TTGTGTNACCGGAGCCAGCGGAGCCATCGGTTTCATGGGTGGTTCGCCTCCTCCTCGAACG : 65

      *           140           *           160           *           180
TrDFRa1 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 180
TrDFRa2 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 180
TrDFRa3 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 180
TrDFRa4 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 180
TrDFRa5 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 178
TrDFRa6 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 178
TrDFRa7 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 175
TrDFRa8 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 175
TrDFRa9 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 174
TrDFRa10 : CGGCTACATTGTCCACGCCACCATCCAAGATCTCGAGGATGAGAACGAGACAAAACATTT : 125

      *           200           *           220           *           240
TrDFRa1 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 240
TrDFRa2 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 240
TrDFRa3 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 240
TrDFRa4 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 240
TrDFRa5 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 238
TrDFRa6 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 238
TrDFRa7 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 235
TrDFRa8 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 235
TrDFRa9 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 234
TrDFRa10 : GGAAGCAATGGAAGGAGCAAAGGGTCATCTCAAATTTTTCGAAATGGATCTTCTTAACAG : 185

```

FIGURE 46

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```

          *           260           *           280           *           300
TrDFRa1 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATNTTGNCTGTCCTAC : 300
TrDFRa2 : TGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATNTTGCATGTGCTAA : 300
TrDFRa3 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAA : 300
TrDFRa4 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAG : 300
TrDFRa5 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAA : 298
TrDFRa6 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAG : 298
TrDFRa7 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAA : 295
TrDFRa8 : TGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTCTTAA : 295
TrDFRa9 : CGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATCTTGCATGTGCTAA : 294
TrDFRa10 : NGACTCTATTGCGGCGCCCGTGAAAGGTTGTGCGCGGAGTTATACATN----- : 232

```

```

          *           320           *           340           *           360
TrDFRa1 : CCGCTTGGGGGANGAGGNNNGCAGCCCN----- : 327
TrDFRa2 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 349
TrDFRa3 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 360
TrDFRa4 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 360
TrDFRa5 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 358
TrDFRa6 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 358
TrDFRa7 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 355
TrDFRa8 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 355
TrDFRa9 : CATCATTTGGTGAAGTCAAAAGACCCCGAGAAGCAAATTTTGAACCGGCAATTCAAGGAAC : 328
TrDFRa10 : ----- : -

```

```

          *           380           *           400           *           420
TrDFRa1 : ----- : -
TrDFRa2 : ----- : -
TrDFRa3 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 420
TrDFRa4 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 420
TrDFRa5 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 418
TrDFRa6 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 418
TrDFRa7 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 415
TrDFRa8 : GGTTAATGTGTTGAAGGTGGCAAAGGAAGCAGGGGTGGAGCGTGTGGTGGCGACATCGTC : 415
TrDFRa9 : ----- : -
TrDFRa10 : ----- : -

```

```

          *           440           *           460           *           480
TrDFRa1 : ----- : -
TrDFRa2 : ----- : -
TrDFRa3 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 480
TrDFRa4 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 480
TrDFRa5 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 478
TrDFRa6 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 478
TrDFRa7 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 475
TrDFRa8 : GATCTCCGCCATCATACCGAGTCCTAATTGGCCAGCTGATAAGATTAAGGGGAGAAGATTG : 475
TrDFRa9 : ----- : -
TrDFRa10 : ----- : -

```

FIGURE 46 (cont)

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```

          *           500           *           520           *           540
TrDFRa1 : ----- : -
TrDFRa2 : ----- : -
TrDFRa3 : NTGGACAGACCTTGATTATTGCAAN500GAAAAAGAAGTTATACTACCCCATTGCAAAGACATT : 540
TrDFRa4 : TTGGACAGACCTTGATTATTGCAAGGAAAAAGAAGTTATACTACCCCATTGCAAAGACACT : 540
TrDFRa5 : TTGGACAGACCTTGATTATTGCAAGGAAAAAGAAGTTATACTACCCCATTGCAAAGACACT : 538
TrDFRa6 : TTGGACAGACCTTGATTATTGCAAGGAAAAAGAAGTTATACTACCCCATTGCAAAGACACT : 538
TrDFRa7 : TTGGACAGACCTTGATTATTGCAAGGAAAAAGAAGTTATACTACCCCATTGCAAAGACATT : 535
TrDFRa8 : TTGGACAGACCTTGATTATTGCAAGGAAAAAGAAGTTA520CTACCCCATTGCAAAGACACT : 535
TrDFRa9 : ----- : -
TrDFRa10 : ----- : -

          *           560           *           580           *           600
TrDFRa1 : ----- : -
TrDFRa2 : ----- : -
TrDFRa3 : AN560CAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGACCGGTTTTGATGTTGTT : 592
TrDFRa4 : AGCAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGACCGGTTTTGATGTTGTTAT : 594
TrDFRa5 : AGCAGAAAAAGCTGGTTGGGAATTTGCTA580AGAGACCGGTTTTG : 583
TrDFRa6 : AGCAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGACCGGTTTTGATGTTGTTATGATTAA : 598
TrDFRa7 : AN560CAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGAC : 571
TrDFRa8 : AGCAGAAAAAGCTGGTTGGGAATTTGCTAAAGAGACCGGTTTTGATGTTGT : 586
TrDFRa9 : ----- : -
TrDFRa10 : ----- : -

          *
TrDFRa1 : ----- : -
TrDFRa2 : ----- : -
TrDFRa3 : ----- : -
TrDFRa4 : ----- : -
TrDFRa5 : ----- : -
TrDFRa6 : CCCTGGTACT : 608
TrDFRa7 : ----- : -
TrDFRa8 : ----- : -
TrDFRa9 : ----- : -
TrDFRa10 : ----- : -

```

FIGURE 46 (cont)

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```

      *           20           *           40           *           60
TrDFRb : ATAAAACCAANCTNCAAAACTGATTGGAAGCTCAGAAAAATAGAAGAAAAGAGATGCCTGA : 60

      *           80           *           100          *           120
TrDFRb : GTTTTGTGTACAGGGGGCACTGGCTTCATAGCAGCCTACCTAGTGAAAGCCTTATTAGA : 120

      *           140          *           160          *           180
TrDFRb : AAAGGGTCACACAGTAAGGACTACTGTAAGAAACCCAGATGATTTGGAGAAGGTTGGTTA : 180

      *           200          *           220          *           240
TrDFRb : TCTAACTGAACTAAGTGAAGACAAAGAGAGATTGAAGATTTTAAAAGCAGATCTATTGGT : 240

      *           260          *           280          *           300
TrDFRb : GGAAGGGAGTTTTGATGAGGCAGTGAGTGGTGTGATGGTGTGTTTCATACAGCTTCCCC : 300

      *           320          *           340          *           360
TrDFRb : TGTTCCTGTTCCACATGATGACAAACATTCAGGTTACTTTGATTGATCCATGTATAAAAGG : 360

      *           380          *           400          *           420
TrDFRb : AACACAAAATGTGCTTAACTCATGCATCAAAGCAAAGGTGAAACGTGTGGTGTTAACATC : 420

      *           440          *           460          *           480
TrDFRb : TTCATGCTCTTCCATAAGATACCGTGACGATGTGCAACAAATTTCTCCTCTTAATGAATC : 480

      *           500          *           520          *           540
TrDFRb : TCATTGGAGTGATTCTGAATACTGCAAACGCTATAACCTGTGGTATGCATATGCAAAGAC : 540

      *           560          *           580          *           600
TrDFRb : TTTAGGAGAAAAAGAAGCATGGAGGATTGCAAAGGAAAGTGGAATTGATCTAGTTGTAGT : 600

      *           620
TrDFRb : TAACCCCTCTTTTGTGGTGGTC : 623

```

## FIGURE 47

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TrDFRb : MPEFCVTGGTGFIAAYLVKALLEKGHTVVRTTVRNPDDLEKVGYLTELSEDKERLKILKAD : 60

TrDFRb : LLVEGSFDEAVSGVDGVFHTASPVLPVPHDDNIQVTLIDPCIKGTQNVNLNSCIKAKVKRVV : 120

TrDFRb : LTSSCSSIRYRDDVQQISPLNESHWS DSEYCKRYNLWYAYAKTLGEKEAWRIAKESGIDL : 180

TrDFRb : VVVNPSFVGG : 190

## FIGURE 48

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```

      *           20           *           40           *           60
TrDFRb1: ATAAAACCAANCTNCAAAACTGATTGGAACCTAGAAAAATAGAAGAAAAGAGATGCCTGA : 60
TrDFRb2: -----GNACTCTCNAAAAATAGAAGAAAAGAGATGCCTGA : 35
TrDFRb3: -----GAAAAGAGATGCCTGA : 16
TrDFRb4: -----GAGATGCCTGA : 11

      *           80           *           100          *           120
TrDFRb1: GTTTTGTGTTACAGGGGGGCACTGGCTTCATAGCAGCCTACCTAGTGAAGCCCTTATTAGA : 120
TrDFRb2: GTTTTGTGTTACAGGGGGGCACTGGCTTCATAGCAGCCTACCTAGTGAAGCCCTTATTAGA : 95
TrDFRb3: GTTTTGTGTTACAGGGGGGCACTGGCTTCATAGCAGCTTACCTAGTGAAGCCCTTATTAGA : 76
TrDFRb4: GTTTTGTGTTACAGGGGGGCACTGGCTTCNTAGCAGCTTACCTAGTGAAGCCCTTATTAGA : 71

      *           140          *           160          *           180
TrDFRb1: AAAGGGTCACACAGTAAGGACTACTGTAAGAAACCCAGATGATTTGGAGAAGGTTGGTTA : 180
TrDFRb2: AAAGGGTCACACAGTAAGGACTACTGTAAGAAACCCAGATGATTTGGAGAAGGTTGGTTA : 155
TrDFRb3: AAAGGGTCACACAGTAAGGACTACTGTAAGAAACCCAGATGATTTGGAGAAGGTTGGTTA : 136
TrDFRb4: AAAGGGTCACACAGTAAGGACTACTGTAAGAAACCCAGATGATTTGGAGAAGGTTGGTTA : 131

      *           200          *           220          *           240
TrDFRb1: TCTAACTGAACCTAAGTGAAGACAAAGAGAGATTGAAGATTTTAAAAGCAGATCTATTGGT : 240
TrDFRb2: TCTAACTGAACCTAAGTGAAGACAAAGAGAGATTGAAGATTTTAAAAGCAGATCTATTGGT : 215
TrDFRb3: TCTAACTGAACCTAAGTGAAGACAAAGAGAGATTGAAGATTTTAAAAGCAGATCTATTGGT : 196
TrDFRb4: TCTAACTGAACCTAAGTGAAGACAAAGAGAGATTGAAGATTTTAAAAGCAGATCTATTGGT : 191

      *           260          *           280          *           300
TrDFRb1: GGAAGGGAGTTTTTGATGAGGCAGTGAGTGGTGTGTTGATGGTGTGTTTCATACAGCTTCCCC : 300
TrDFRb2: GGAAGGGAGTTTTTGATGAGGCAGTGAGTGGTGTGTTGATGGTGTGTTTCATACAGCTTCCCC : 275
TrDFRb3: GGAAGGGAGTTTTTGATGAGGCAGTGAGTGGTGTGTTGATGGTGTGTTTCATACAGCTTCCCC : 256
TrDFRb4: GGAAGGGAGTTTTTGATGAGGCAGTGAGTGGTGTGTTGATGGTGTGTTTCATACAGCTTCCCC : 251

      *           320          *           340          *           360
TrDFRb1: TGTTCCTTGTTCACATGATGACAAATTTCAGGTTACTTTTGATTGATCCATGTATAAAAGG : 360
TrDFRb2: TGTTCCTTGTTCACATGATGACAAATTTCAGGTTACTTTTGATTGATCCATGTATAAAAGG : 335
TrDFRb3: TGTTCCTTGTTCACATGATGACAACATTTCAGGTTACTTTTGATTGATCCATGTATAAAAGG : 316
TrDFRb4: TGTTCCTTGTTCACATGATGACAACATTTCAGGTTACTTTTGATTGATCCATGTATAAAAGG : 311

      *           380          *           400          *           420
TrDFRb1: AACACAAAATGTGCTTAACCTCATGCATCAAAGCAAAGGTGAAACGTGTGGTGTTAACATC : 420
TrDFRb2: AACACAAAATGTGCTTAACCTCATGCATCAAAGCAAAGGTGAAACGTGTGGTGTTAACATC : 395
TrDFRb3: AACACAAAATGTGCTTAACCTCATGCATCAAAGCAAAGGTGAAACGTGTGGTGTTAACATC : 376
TrDFRb4: AACACAAAATGTGCTTAACCTCATGCATCAAAGCAAAGGTGAAACGTGTGGTGTTAACATC : 371

      *           440          *           460          *           480
TrDFRb1: TTCATGCTCTTCCATAAGATACCGTGACGATGTGCAACAAATTTCTCCTCTTAATGAATC : 480
TrDFRb2: TTCATGCTCTTCCATAAGATACCGTGACGATGTGCAACAAATTTCTCCTCTTAATGAATC : 455
TrDFRb3: TTCATGCTCTTCCATAAGATACCGTGACGATGTGCAACAAATTTCTCCTCTTAATGAATC : 436
TrDFRb4: TTCATGCTCTTCCATAAGATACCGTGACGATGTGCAACAAATTTCTCCTCTTAATGAATC : 431

```

FIGURE 49



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```

          *           500           *           520           *           540
TrDFRb1: TCATTGGAGTGATTCTGAATACTGCAAACGCTATAACCTGTGGTATGCATATGCAAAGAC : 540
TrDFRb2: TCATTGGAGTGATTCTGAAT TACTGCAAACGCTATAACCTGTGGTATGCATATGCAAAGAC : 515
TrDFRb3: TCATTGGAGTGATTCTGAATACTGCAAACGCTATAACCTGTGGTATGCATATGCAAAGAC : 496
TrDFRb4: TCATTGGAGTGATTCTGAATACTGCAAACGCTATAACCTGTGGTATGCATATGCAAAGAC : 491

```

```

          *           560           *           580           *           600
TrDFRb1: TTTAGGAGAAAAAGAAGCATGGAGGATTGC----- : 570
TrDFRb2: TTTAGGAGAAAAAGAAGCATGGAGGATTGCAAAGGAAAGTGGGATTAATCTAGTTGT--- : 572
TrDFRb3: TTTAGGAGAAAAAGAAGCATGGAGGATTGCAAAGGAAAGTGGGAATTGATCTAGTTGTAGT : 556
TrDFRb4: TTTAGGAGAAAAAGAAGCATGGAGGATTGCAAAGGAAAGTGGGAATTGATCTAGTTGTAGT : 551

```

```

          *           620
TrDFRb1 : ----- : -
TrDFRb2 : ----- : -
TrDFRb3 : TAACCCCTCTTTTGTT----- : 572
TrDFRb4 : TAACCCCTCTTTTGTTGGTGTC : 574

```

FIGURE 49 (cont)

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```

      *           20           *           40           *           60
TrDFRc : GNGAAGANCTAGTTTGCCTAACCCGGANCAACGGTTTCATCGGAACATGGCTAGTTAAAA : 60

      *           80           *           100          *           120
TrDFRc : CCCTACTCCAAAAACACTACAAAATTACGCCACAATCTTCCCCAATTCCAACGCATCTC : 120

      *           140          *           160          *           180
TrDFRc : ATCTCTTCACACTCCACCCGGAAGCTCAATCCCGGATCACAATTTCCCTGTCGATATCC : 180

      *           200          *           220          *           240
TrDFRc : TCGACTCCACCGCCGTCTTCTCCGCTATCAATAACTGCTCAGGTGTCTTTCATGCCGCTT : 240

      *           260          *           280          *           300
TrDFRc : CTCCATGTACCCTCGAAGATCCAACCTGATCCGCAAAAAGAGCTTCTAGAACCTGCTGTAC : 300

      *           320          *           340          *           360
TrDFRc : AAGGAACCCTAAATGTTCTAGAAAGCATCCAGCGCGCAGGTACCAAACCCTAATTGGCCGG : 360

      *           380          *           400          *           420
TrDFRc : AGAAAAAGGCGATCGATGAGGCGTCGTGGACGGATGTTGAGTACTGTAAATTGAGAGGGA : 420

      *           440          *           460          *           480
TrDFRc : AGTGGTATCTGGTGTGCAAAACGGAGGCGGAGAAGGCGGCTTGGGATTTTCGAGAGAAAA : 480

      *           500          *           520          *           540
TrDFRc : ATGGTGGTGTGTTGATGTGGGGGCGGNTCATCCGGGGACTTGTTTGGGAGAGTTGATACAGA : 540

      *           560          *           580          *           600
TrDFRc : AGGAGTTGAATGCGAGTTACAGCGGNTTTACAGAGGTTGATGATGGGGAGTGAGGATACTC : 600

      *           620          *           640
TrDFRc : AAGAGTGNTATTGGNNGGGGGGCTGNNNATGNTAAAGATGN : 641

```

## FIGURE 50

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TrDFRc :           \*          20          \*          40          \*          60  
          EXLVCVTRXNGFIGTWLVKTLQKHYKIHATIFPNSNASHLFTLHPEAQSRITIFPVDIL : 60

TrDFRc :           \*          80          \*          100          \*          120  
          DSTAVFSAINNCSGVFHAASPCTLEDPTDPQKELLEPAVQGTNLVLEASSAQVPNPWPPE : 120

TrDFRc :           \*          140          \*          160          \*          180  
          KKAIDEASWTDVEYCKLRGKWYLVSKTEAEKAAWDFREKNGGVDVGAXHPGTCLGELIQK : 180

TrDFRc :           \*          200  
          ELNASSALQRLMMGSEDTEQEXYWXGG : 206

## FIGURE 51

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```

      *           20           *           40           *           60
TrDFRd : GCNTTGATCAGCGTATNAACACACANGTCTTCCCTTGAGCTCTGTTTCTCCACATGTCGA : 60

      *           80           *           100          *           120
TrDFRd : AGCTAGTTTGCGTCACCGGCGGCAGCGGATGCATCGGTTTCATGGCTAGTCCATCTCCTTC : 120

      *           140          *           160           *           180
TrDFRd : TCCTCCGCGGCTACACTGTTTCAGCCACCGTCCAAAATCTCAATGATGAGAACGAAACGA : 180

      *           200          *           220           *           240
TrDFRd : AGCATCTAGAAGCTCTCGAAGGAGCACAACTAATCTCCGTCTCTTCCAGATCGATCTCC : 240

      *           260          *           280           *           300
TrDFRd : TTAAC TACGACACAATCCTCGCTGCTGTCCGCGGTTGCGTCGGAATTTTCCACCTCGCTT : 300

      *           320          *           340           *           360
TrDFRd : CACCTTGCACTGTAGACAAAGTTCATGATCCTCAGAAGGAGCTTTTGGATCCTGCAATTA : 360

      *           380          *           400           *           420
TrDFRd : AAGGGACTTTGAATGTGCTTACTGCAGCTAAGGAAGTAGGGGTGAAGCGTGTGGTTGTTA : 420

      *           440          *           460           *           480
TrDFRd : CCTCGTCTGTCTCGGCGATTACTCCTAGTCTGATTGGCCTTCTGATGTTGTTAAAAGAG : 480

      *           500          *           520           *           540
TrDFRd : AGGATTGTTGGACTGATGTTGAATATTGCAAGAAAAAAGAGTTGGGGTATCCGTTGTCCA : 540

      *           560          *           580           *           600
TrDFRd : AAACATTGGCTGAAAAAGCTGCGTGGGATTTTTNCAAAGAAAATGTTTGGATGTTGTTG : 600

      *           620          *           640           *           660
TrDFRd : NGGTGAATCCCGNACTGNGATGGGTCCTGTTTTTCCACCACGGCATAATGCAAGCATGC : 660

      *           680          *           700           *           720
TrDFRd : TCATGCCTTGGGAACTTTTTTGAAGGCTGGNNCTGAAACATTTGAAGACTATTTTATGG : 720

      *           740          *           760           *           780
TrDFRd : GATTGGCCNNCTTTAAAGATGTNGCATTGGCNCATNNTTTGGGGTATGAGAACAAANANN : 780

      *           800          *           820
TrDFRd : CTTTGGGANACATNGGNGGGTTGAAACTATCNNTCCTTACGG : 822

```

## FIGURE 52

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TrDFRd : MSKLVCVTGGSGCIGSWLVHLLLLRGYTVHATVQNLNDENETKHLEALEGAQTNLRRLFQI : 60

TrDFRd : DLLNYDTILAAVRGCVGIFHLASPCTVDKVHDPQKELLDPAIKGTNLNVLTAKEVGVKRV : 120

TrDFRd : VVTSSVSAITPSPDWPSDVVKREDCWTDVEYCKKKELGYPLSKTLAEKAAWDFXKENGLD : 180

TrDFRd : VVXVNPXTXMGPFVFPFRHNASMLMPWETFLKAXXETFEDYFMGLAXFKDXALXHXLYEN : 240

TrDFRd : KXXLGXXXGLKLXXLT : 256

## FIGURE 53

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	*                      20                      *                      40                      *                      60	
TrDFRd1:	GCNTTGATCAGCGTATNAACACACANGTCTTCCCTTGAGCTCTGTTTCTCCNCATGTCGA	: 60
TrDFRd2:	-----MACANNTCTTCCCTTGAGCTCTGTTTCTCCACATGTCGA	: 39
	*                      80                      *                      100                      *                      120	
TrDFRd1:	AGCTAGTTTGCGTCACCGGCGGCAGCGGATGCATCGGTTTCATGGCTAGTCCATCTCCTTC	: 120
TrDFRd2:	AGCTAGTTTGCGTCACCGGCGGCTGCGGATGCATCGGTTTCATGGCTAGTCCATCTCCTTC	: 99
	*                      140                      *                      160                      *                      180	
TrDFRd1:	TCCTCCGCGGCTACACTGTTTACGCCACCGTCCAAAATCTCAATGATGAGAACGAAACGA	: 180
TrDFRd2:	TCCTCCGCGGCTACACTGTTTACGCCACCGTCCAAAATCTCAATGATGAGAACGAAACGA	: 159
	*                      200                      *                      220                      *                      240	
TrDFRd1:	AGCATCTAGAAGCTCTCGAAGGAGCACAACTAATCTCCGTCTCTTCCAGATCGATCTCC	: 240
TrDFRd2:	AGCATCTAGAAGCTCTCGAAGGAGCACAACTAATCTCCGTCTCTTCCAGATCGATCTCC	: 219
	*                      260                      *                      280                      *                      300	
TrDFRd1:	TTAACTACGACACAATCCTCGCTGCTGTCCGCGGTTGCGTCGGAATTTCCACCTCGCTT	: 300
TrDFRd2:	TTAACTACGACACAATCCTCGCTGCTGTCCGCGGTTGCGTCGGAATTTTCCACCTCGCTT	: 279
	*                      320                      *                      340                      *                      360	
TrDFRd1:	CACCTTGCACTGTAGACAAAAGTTCATGATCCTCAGAAGGAGCTTTTGGATCCTGCAATTA	: 360
TrDFRd2:	CACCTTGCACTGTAGACAAAAGTTCATGATCCTCAGAAGGAGCTTTTGGATCCTGCAATTA	: 339
	*                      380                      *                      400                      *                      420	
TrDFRd1:	AAGGGACTTTGAATGTGCTTACTGTCAGCTAAGGAAGTAGGGGTGAAGCGTGTGGTTGTTA	: 420
TrDFRd2:	AAGGGACTTTGAATGTGCTTACTGTCAGCTAAGGAAGTAGGGGTGAAGCGTGTGGTTGTTA	: 399
	*                      440                      *                      460                      *                      480	
TrDFRd1:	CCTCGTCTGTCTCGGCGATTACTCCTAGTCCTGATTGGCCTTCTGATGTTGTTAAAAGAG	: 480
TrDFRd2:	CCTCGTCTGTCTCGGCGATTACTCCTAGTCCTGATTGGCCTTCTGATGTTGTTAAAAGAG	: 459
	*                      500                      *                      520                      *                      540	
TrDFRd1:	AGGATTGTTGGACTGATGTTGAATATTGCAAGAAAAAGAGTTCTGGGTATCCGTT-----	: 535
TrDFRd2:	AGGATTGTTGGACTGATGTTGAATATTGCAAGAAAAAGAGTTGGGGTATCCGTTGTCCA	: 519
	*                      560                      *                      580                      *                      600	
TrDFRd1:	-----	: -
TrDFRd2:	AAACATTGGCTGAAAAAGCTGCGTGGGATTTTTNCAAGAAAATGGTTTGGATGTTGTTG	: 579
	*                      620                      *                      640                      *                      660	
TrDFRd1:	-----	: -
TrDFRd2:	NGGTGAATCCCGNACTGNGATGGGTCTGTTTTTCCACCACGGCATAATGCAAGCATGC	: 639

FIGURE 54

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TrDFRd1:	-----*	680	*	700	*	720	-----	:	-	
TrDFRd2:	TCATGCCTTGGGAAACTTTTTTGAAGGCTGGNNCTGAAACATTGAAGACTATTTTATGG								:	699
TrDFRd1:	-----*	740	*	760	*	780	-----	:	-	
TrDFRd2:	GATTGGCCNNCTTTAAAGATGTNGCATTGGCNCATNNTTTGGGGTATGAGAACAAANANN								:	759
TrDFRd1:	-----*	800	*	820	-----			:	-	
TrDFRd2:	CTTTGGGANACATNGGNGGGTTGAAACTATCNNTCCTTACGG								:	801

**FIGURE 54 (cont)**

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      *           20           *           40           *           60
TrDFRe : GTCAC TTATGAAATGGAACACAAAGGTGGAGACAAAGTATGTGTGACAGGGGCATCAGGC : 60

      *           80           *           100          *           120
TrDFRe : TTTT TAGCATCTTGGCTTATTAAGAACTTCTTTTGTCTGGCTATCAAGTCATTGGAACA : 120

      *           140          *           160          *           180
TrDFRe : GTTAGAGATTTAGGGAAGAAGAAGAAAGTTGAACATTTATGGAAATTGGAAGGAGCAACA : 180

      *           200          *           220          *           240
TrDFRe : GAAAGACTAGAACTAATCCAAGCTGATTTAATGGAAGAAAATAGTTTCGACAAAGCGATC : 240

      *           260          *           280          *           300
TrDFRe : ATGGGATGCAAAGGTGTCTTCCACATTGCCTCTCCAGTACTCAATCATATATCAGATAAT : 300

      *           320          *           340          *           360
TrDFRe : CCTAAGGCGGAAATCTTGGAACCGGCAGTCCAAGGTACGCTAAATGTGTTGCGTTCTTGT : 360

      *           380          *           400          *           420
TrDFRe : AAGAGGAACCCCGATCTTGTTTCGAGTGGTGCTAGCCTCATCATCTTCGGCTGTTAGAGTA : 420

      *           440          *           460          *           480
TrDFRe : AGAGCTGATTTTGATCCAAGCATACCAATTGATGAATCATCTTGGAGCTCCTTGGAATTG : 480

      *           500          *           520          *           540
TrDFRe : TGCGAGAAACTCAAGGCATGGTACCCAATGTCAAAGACAATGGCAGAAAAAGCAGCTTGG : 540

      *           560          *           580
TrDFRe : GAATATAGCAAAGAGAATGGAATAGACTTAGTGACTATTTTC : 582

```

## FIGURE 55



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TrDFRe : MEHKGGDKVCVTGASGFLASWLIKLLLSGYQVIGTVRDLGKKKKVEHLWKLEGATERLE : 60

TrDFRe : LIQADLMEENSFDKAIMGCKGVFHIA SPVLNHISDNPKAEILEPAVQGTNLNVLRSCKRNP : 120

TrDFRe : DLVRVVLASSSSAVRVRADFDPSIPIDESSWSSLELCEKLKAWYPMSKTMAEKA AWEYSK : 180

TrDFRe : ENGIDLVTIF : 190

## FIGURE 56

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```

      *           20           *           40           *           60
TrDFRf : TNCNNGCTNCNTNCGGGCAGAGANTTTCCTGACCTATNTGTTACTNAAGAATATTTCTA : 60

      *           80           *           100          *           120
TrDFRf : TATATATATTTGTGTTTCAAGAACCCAAAAAATAGAATAGTGATGGAAAGGAGTTGCAAG : 120

      *           140          *           160          *           180
TrDFRf : GTTTGTGTCACCGGTGGTGTGTTATATTGGTTCTCTTTTAGTCAAAAAGCTTTTGGAA : 180

      *           200          *           220          *           240
TrDFRf : AAGGGTTACACCGTTCATGCTACTCTTAGAACTTGAAGGACGAATCCAAAGTAGATTTT : 240

      *           260          *           280          *           300
TrDFRf : TTGAGAGGCTTCCACATGCAGATACTAGACTTATGTTATTTGAAGCTGATATATACAAA : 300

      *           320          *           340          *           360
TrDFRf : TCAGATGAATTTTGGCCCGCAATTCAAGGTTGTGAGTTTGTTTTTCACCTTGCTACTCCT : 360

      *           380          *           400          *           420
TrDFRf : TTTCAACATCAAACCTGATTCTCAGTTTAAGAGCATAGAGGAAGCTGCAATAGCAGGGGTA : 420

      *           440          *           460          *           480
TrDFRf : AAAAGCATAGCTGAAAATTGCATAAAATCAGGAACAGTGAGAAAATTGATATACACTGGA : 480

      *           500          *           520          *           540
TrDFRf : ACTGTAATTGCTTCTTCTTCTCTGAAAGATGATGGAAGTGGCTACAAAGACTTCATTGAT : 540

      *           560          *
TrDFRf : GAAACTTGTTGGACACCTCTCCATCTTCCTCT : 572

```

## FIGURE 57

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TrDFRf : MERSCKVCVTGGAGYIGSLLVKKLLEKGYTVHATLRNLKDESKVDFLRGFPHADTRLMLF : 60

TrDFRf : EADIYKSDEFWPAIQGCEFVFHLATPFQHQTD SQFKSIEEAAIAGVKSAENCIKSGTVR : 120

TrDFRf : KLIYTGTVIASSSLKDDGSGYKDFIDETCWTPHLHP : 156

**FIGURE 58**

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TrDFRg :                   \*                  20                  \*                  40                  \*                  60  
GCCNTTGCCTACTACTAAACTATATATTATTATTATTATATATGATGATACATAGTGACA : 60

TrDFRg :                   \*                  80                  \*                  100                  \*                  120  
TTAATAATTGGAAGGGAGAATAAATAGTTGAAAAACACACAGTTGGAGTGTTTTTGTGT : 120

TrDFRg :                   \*                  140                  \*                  160                  \*                  180  
TAAAGAAGCTNGAAAATGGAGGAAGCNACAAAGATGGTGAAAAAGAGTGGACAAATTGTT : 180

TrDFRg :                   \*                  200                  \*                  220                  \*                  240  
CCTACTGCCAAATACTGTGTTACAGGAGCAACAGGCTATATTGGTTCATGGCTTGTTGAA : 240

TrDFRg :                   \*                  260                  \*                  280                  \*                  300  
GCTCTTCTTCAAAGAGGTTGCACTGTTTCATGCTACTGTTAGAGATCCTGAAAAATCGTTA : 300

TrDFRg :                   \*                  320                  \*                  340                  \*                  360  
CACCTCCTGTCTGTTGTGGAAAGGTAGTGACCAATTGAGAATTTTCCGTGCGGATTTGCAA : 360

TrDFRg :                   \*                  380                  \*                  400                  \*                  420  
GAAGAAGGAAGTTTCGATGATGCCGTAAAAGGATGTATTGGTGTGTTCCATGTTGCAGCT : 420

TrDFRg :                   \*                  440                  \*                  460                  \*                  480  
TCAATGCAATTCAATATTAGTGACAAAGAAAACACTGAGGACTTTGTTGAAGCAAATATA : 480

TrDFRg :                   \*                  500                  \*                  520                  \*                  540  
ATTGACCCTGCAATCAAAGGAACCATAAATCTTCTCAAATCATGCTTGAAATCAAATTCA : 540

TrDFRg :                   \*                  560                  \*                  580                  \*                  600  
GTGAAAAGGGTTGTTTTACATCTTCCATAAGTACTATTACTGCTAAAGACAACGACGGA : 600

TrDFRg :                   \*                  620                  \*                  640                  \*                  660  
AAATGGAAACCTATTGTTGATGAATCTTGCCAAACAAAACCTGAGATTCTGTGGAATACA : 660

TrDFRg :                   \*                  680                  \*                  700                  \*                  714  
CAACCAAGTGGATGGGTTTATGCACTTTCAAAGCTTCATGCAGAAGAAGCGGCT : 714

## FIGURE 59

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TrDFRg : MVKKSGQIVPTAKYCVTGATGYIGSWLVEALLQRGCTVHATVRDPEKSLHLLSLWKGSDQ : 60

TrDFRg : LRIFRADLQEEGSFDDAVKGCIGVFHVAASMQFNISDKENTEDFVEANIIDPAIKGTINL : 120

TrDFRg : LKSCCLKSNSVKRVVFTSSISTITAKDNDGKWKPIVDESCQTKTEILWNTQPSGWVYALSK : 180

TrDFRg : LHAEEAA : 187

## FIGURE 60

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	*                      20                      *                      40                      *                      60	
TrDFRg1:	<b>GCCNTTGCCTACTACTAAACTATATATTATTATTATTATATATGATGATACATAGTGACA</b>	: 60
TrDFRg2:	-----	: -
TrDFRg3:	-----	: -
	*                      80                      *                      100                      *                      120	
TrDFRg1:	<b>TTAATAATTGGAAGGGAGAAATAAATAGTTGAAAAACACACAGTTGGAGTGTTTTTGTTGT</b>	: 120
TrDFRg2:	-----	: -
TrDFRg3:	-----	: -
	*                      140                      *                      160                      *                      180	
TrDFRg1:	<b>TAAAGAAGCTTAAAAATGGAGGAAACACAAAGATGGTGAAAAAAGTGGACAAATTGTT</b>	: 180
TrDFRg2:	---AGAAGCTNGAAAAATGGAGGAAGACACAAAGATGGTGAAAAAAGTGGACAAATTGTT	: 57
TrDFRg3:	-----CTNGAAAAATGGAGGAAGCNACAAAGATGGTGAAAAAAGTGGACAAATTGTT	: 52
	*                      200                      *                      220                      *                      240	
TrDFRg1:	<b>CCTAAGCCAAATACTGTGTACAGGAGCACAGGCTATATTGGTTCATGGCTTGTTGAA</b>	: 240
TrDFRg2:	<b>CCTACTGCCAAATACTGTGTACAGGAGCAACAGGCTATATTGGTTCATGGCTTGTTGAA</b>	: 117
TrDFRg3:	<b>CCTACTGCCAAATACTGTGTACAGGAGCAACAGGCTATATTGGTTCATGGCTTGTTGAA</b>	: 112
	*                      260                      *                      280                      *                      300	
TrDFRg1:	<b>GCTCTTCTTCAAAGAGGTTGCAGTGTTCATGCTACTGTTAGAGATCCTG</b> -----	: 289
TrDFRg2:	<b>GCTCTTCTTCAAAGAGGTTGCAGTGTTCATGCTACTGTTAGAGATCCTGAAAAATCGTTA</b>	: 177
TrDFRg3:	<b>GCTCTTCTTCAAAGAGGTTGCAGTGTTCATGCTACTGTTAGAGATCCTGAAAAATCGTTA</b>	: 172
	*                      320                      *                      340                      *                      360	
TrDFRg1:	-----	: -
TrDFRg2:	<b>CACCTCCTGTCGTTGTGGAAGGTAGTGACCAATTGAGAATTTCCGTGCGGATTTGCAA</b>	: 237
TrDFRg3:	<b>CACCTCCTGTCGTTGTGGAAGGTAGTGACCAATTGAGAATTTCCGTGCGGATTTGCAA</b>	: 232
	*                      380                      *                      400                      *                      420	
TrDFRg1:	-----	: -
TrDFRg2:	<b>GAAGAAGGAAGTTTCGATGATGCCGTAAAAGGATGTATTGGTGTGTTCCATGTTGCAGCT</b>	: 297
TrDFRg3:	<b>GAAGAAGGAAGTTTCGATGATGCCGTAAAAGGATGTATTGGTGTGTTCCATGTTGCAGCT</b>	: 292
	*                      440                      *                      460                      *                      480	
TrDFRg1:	-----	: -
TrDFRg2:	<b>TCAATGCAATTCAATATTAGTGACAAAGAAAACACTGAGGACTTTGTTGAAGCAAATATA</b>	: 357
TrDFRg3:	<b>TCAATGCAATTCAATATTAGTGACAAAGAAAACACTGAGGACTTTGTTGAAGCAAATATA</b>	: 352
	*                      500                      *                      520                      *                      540	
TrDFRg1:	-----	: -
TrDFRg2:	<b>ATTGACCCTGCAATCAAAGGAACCATAAATCTTCTCAAATCATGCTTGAAATCAAATTCA</b>	: 417
TrDFRg3:	<b>ATTGACCCTGCAATCAAAGGAACCATAAATCTTCTCAAATCATGCTTGAAATCAAATTCA</b>	: 412

FIGURE 61

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TrDFRg1:	-----*	560	*	580	*	600	-----	:	-		
TrDFRg2:	GTGAAAAGGGTTGTTTTTCACATCTTCCATAAGTACTATTACTGCTAAAGACAACGACGGA									:	477
TrDFRg3:	GTGAAAAGGGTTGTTTTTCACATCTTCCATAAGTACTATTACTGCTAAAGACAACGACGGA									:	472
TrDFRg1:	-----*	620	*	640	*	660	-----	:	-		
TrDFRg2:	AAATGGAAACCTATTGTTGATGAATCTTGCCAAACAAAACTGAGATTCTGTGGAATACA									:	537
TrDFRg3:	AAATGGAAACCTATTGTTGATGAATCTTGCCAAACAAAACTGAGATTCTGTGGAATACA									:	532
TrDFRg1:	-----*	680	*	700	*		-----	:	-		
TrDFRg2:	CAACCAAGTGGATGGGTTTATGCACTTTCAAAGCTTCATGCAGAAGAAGCGGCT									:	591
TrDFRg3:	CAACCAAGTGGATGGGTTTATGCACTTTCAAAGCTTCATG-----									:	572

**FIGURE 61 (cont)**

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TrDFRh : GNNNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAGGAAAAAAGAGTAGAGAAGTGA : 60  
 \* 20 \* 40 \* 60

TrDFRh : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 120  
 \* 80 \* 100 \* 120

TrDFRh : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 180  
 \* 140 \* 160 \* 180

TrDFRh : ACCGTTTCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA : 240  
 \* 200 \* 220 \* 240

TrDFRh : AAAAGCAAATTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA : 300  
 \* 260 \* 280 \* 300

TrDFRh : ATTAAAGGGTGACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC : 360  
 \* 320 \* 340 \* 360

TrDFRh : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 420  
 \* 380 \* 400 \* 420

TrDFRh : GAAAAGGCAAAAACAGTTAGAAAATTGGTTTTACATCATCGGCTGGAAGTGTGGACGTT : 480  
 \* 440 \* 460 \* 480

TrDFRh : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGCCGT : 540  
 \* 500 \* 520 \* 540

TrDFRh : AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT : 600  
 \* 560 \* 580 \* 600

TrDFRh : TGGAAGTATTCGAAAGAGCACAAACATAGACTTTGTCTCCATCATTCCACCTCTTGTGTT : 660  
 \* 620 \* 640 \* 660

TrDFRh : GGCCCCCTTCTTATGGCCTCAATGCCACCTAGTCTAATCACTGCTCTTTCTCTTATCACA : 720  
 \* 680 \* 700 \* 720

TrDFRh : GGAAATGAGGCCCATTAATCAATCATAAAGCAAGGGCAATACGTCCATTTAGATGACCTT : 780  
 \* 740 \* 760 \* 780

TrDFRh : TGTCTTGCTCATATATTTCTGTATGAGAATCCAAAAGCTCAAGGGAGATACATTTGCTGT : 840  
 \* 800 \* 820 \* 840

## FIGURE 62



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TrDFRh : TCACATGAAGCA<sup>\*</sup>ACCATTCATCAAGTTGCAAACTTATTAAAGAAAAATACCCAGAGTTC : 900  
860 880 900

TrDFRh : AATGTCCCAACAAAATTCAATGATATCCAGATGAATTGGAAATTATTAAATTTTCTAAA : 960  
920 940 960

TrDFRh : AAGAAGATCACAGACT : 976  
976

**FIGURE 62 (cont)**

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TrDFRh : MGSESEIVCVTGASGFIGSWLVMRLIERGYTVRATVRDPDNMKKVKHLVELPGA KSKLSL : 60

TrDFRh : WKADLDKEGSFDEAIKGCTGVFHVATPMDFESKDPENEVIKPTINGLIDILKACEKAKTV : 120

TrDFRh : RKL VFTSSAGTVDVTEHPKSIIDETCWS D VDFCRRVKMTGWMYFVSKTLAEQEAWKYSKE : 180

TrDFRh : HNIDFVSIIPPLVVGPFLMASMPPSLITALSLITGNEAHYSIIKQGQYVHLDDLCLAHIF : 240

TrDFRh : LYENPKAQGRYICCSHEATIHQVAKLIKEKYPEFNVPTKFNDIPDELEIIKFSKKKITD : 299

## FIGURE 63

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      *           20           *           40           *           60
TrDFRh1 : GNN TGTCTTCCGTTNAATTTNAGNCTATATTGAAAAAGAAAAAGAGTACNNNANNGA : 60
TrDFRh2 : --- TGTCTTAAATCAATTTACNCTATATTGAAAAGGAAAAAGAGGAGAGAACTCA : 59
TrDFRh3 : --- GNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAAGAAAAAGAGTAGNNNANNGA : 57
TrDFRh4 : --- GNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAAGAAAAAGAGTAGAGAAANNNGN : 57
TrDFRh5 : --- GNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAGNAAAAAGAGTAGNNNANNGA : 57
TrDFRh6 : --- GNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAGNAAAAAGAGTACNNNANNGA : 57
TrDFRh7 : --- GNGTCTTCCGTTNAATTTNAGNCTATATTGAAAAAGAAAAAGAGTAGAGAACTGA : 57
TrDFRh8 : --- GGTCTTCCATTNAATTTNAGNCTATATTGAAAAGGAAAAAGAGGAGAGAACTGA : 56
TrDFRh9 : --- GGTCTTCCGTTNAATTTNAGNCTATATTGAAAAAGAAAAAGAGTAGAGAACTGA : 56
TrDFRh10 : --- GGTCTTCCGTTNATTTNAGNCTATATTGAAAAAGAAAAAGAGTAGAGAACTGA : 56
TrDFRh11 : --- GTCTTCNNTTNAATTTNAGNCTATATTGAAAAGGAAAAAGAGGAGAGAACTGA : 55
TrDFRh12 : --- GTCTTCCGTTNATTTNAGNCTATATTGAAAAGNAAAAAGAGTAGAGAAANNNGN : 55
TrDFRh13 : --- GTCTTCTTNAATTTCCACTATATTGAAAAGGAAAAAGAGGAGAGAACTGA : 52
TrDFRh14 : ----- GNAGAGAACTNA : 12
TrDFRh15 : ----- : -

      *           80           *           100          *           120
TrDFRh1 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 120
TrDFRh2 : AATGAGTGAAAAATACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 119
TrDFRh3 : AGTGAAGTGAAATACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 117
TrDFRh4 : NNTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 117
TrDFRh5 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 117
TrDFRh6 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 117
TrDFRh7 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 117
TrDFRh8 : AATGAAGTGAAAAATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 116
TrDFRh9 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 116
TrDFRh10 : AGTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 116
TrDFRh11 : AATGAAGTGAAAAATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 115
TrDFRh12 : NNTGAAGTGAAAACATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 115
TrDFRh13 : AATGAGTGAAAAATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 112
TrDFRh14 : CATGAGTGAAAAATACATGGGTTCCGAATCAGAAATAGTTTGTGTTACCGGAGCTTCA : 72
TrDFRh15 : ----- : -

      *           140          *           160          *           180
TrDFRh1 : GTTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 180
TrDFRh2 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 179
TrDFRh3 : GTTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 177
TrDFRh4 : GTTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 177
TrDFRh5 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 177
TrDFRh6 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 177
TrDFRh7 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 177
TrDFRh8 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 176
TrDFRh9 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 176
TrDFRh10 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 176
TrDFRh11 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 175
TrDFRh12 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACGGTTCGAGCC : 175
TrDFRh13 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACTGTTTCGAGCC : 172
TrDFRh14 : GGTTCATCGGATCGTGGCTTGTTATGAGACTTATAGAGCGTGGCTACACTGTTTCGAGCC : 132
TrDFRh15 : ----- : -

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FIGURE 64

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      *           200           *           220           *           240
TrDFRh1 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :240
TrDFRh2 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :239
TrDFRh3 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :237
TrDFRh4 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :237
TrDFRh5 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :237
TrDFRh6 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :237
TrDFRh7 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :237
TrDFRh8 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :236
TrDFRh9 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :236
TrDFRh10 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :236
TrDFRh11 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :235
TrDFRh12 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :235
TrDFRh13 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :232
TrDFRh14 : ACCGTTCCGCGACCCAGATAACATGAAGAAGGTGAAGCATTTGGTGGAACTGCCGGGTGCA :192
TrDFRh15 : ----- : -

      *           260           *           280           *           300
TrDFRh1 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :300
TrDFRh2 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :299
TrDFRh3 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :297
TrDFRh4 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :297
TrDFRh5 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :297
TrDFRh6 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :297
TrDFRh7 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :297
TrDFRh8 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :296
TrDFRh9 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :296
TrDFRh10 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :296
TrDFRh11 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :295
TrDFRh12 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :295
TrDFRh13 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :292
TrDFRh14 : AAAAGCAAATTTGTCTCTTTGGAAGGCTGATCTTGATAAAGAGGGGAGTTTTGATGAAGCA :252
TrDFRh15 : ----- : -

      *           320           *           340           *           360
TrDFRh1 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :360
TrDFRh2 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :359
TrDFRh3 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :357
TrDFRh4 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :357
TrDFRh5 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :357
TrDFRh6 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :357
TrDFRh7 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :357
TrDFRh8 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :356
TrDFRh9 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :356
TrDFRh10 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :356
TrDFRh11 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :355
TrDFRh12 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :355
TrDFRh13 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :352
TrDFRh14 : ATTAAAGGGTGCACAGGAGTTTTTCATGTTGCTACACCAATGGATTTTGAATCCAAGGAC :312
TrDFRh15 : ----- : -

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FIGURE 64 (cont)

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      *           380           *           400           *           420
TrDFRh1 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 420
TrDFRh2 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 419
TrDFRh3 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 417
TrDFRh4 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 417
TrDFRh5 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 417
TrDFRh6 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 417
TrDFRh7 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 417
TrDFRh8 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 416
TrDFRh9 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 416
TrDFRh10 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 416
TrDFRh11 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 415
TrDFRh12 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 415
TrDFRh13 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 412
TrDFRh14 : CCTGAGAATGAAGTGATAAAGCCTACAATAAATGGATTAATAGACATACTGAAAGCATGC : 372
TrDFRh15 : -----TACTGAAAGCATGC : 14

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      *           440           *           460           *           480
TrDFRh1 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 480
TrDFRh2 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 479
TrDFRh3 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 477
TrDFRh4 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 477
TrDFRh5 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 477
TrDFRh6 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 477
TrDFRh7 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 477
TrDFRh8 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 476
TrDFRh9 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 476
TrDFRh10 : GAAAAGGCCAAAAACAGTTAATAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 476
TrDFRh11 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 475
TrDFRh12 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 475
TrDFRh13 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 472
TrDFRh14 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 432
TrDFRh15 : GAAAAGGCCAAAAACAGTTAGAAAATTTGGTTTTTCACATCATCGGCTGGAACGTGTGGACGTT : 74

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      *           500           *           520           *           540
TrDFRh1 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 540
TrDFRh2 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 539
TrDFRh3 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 493
TrDFRh4 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 537
TrDFRh5 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 537
TrDFRh6 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 537
TrDFRh7 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 536
TrDFRh8 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 536
TrDFRh9 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 536
TrDFRh10 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 535
TrDFRh11 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 535
TrDFRh12 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 532
TrDFRh13 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 492
TrDFRh14 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 134
TrDFRh15 : ACTGAACATCCAAAGTCTATTATTGATGAAACATGCTGGAGTGACGTTGACTTTTGGCCGT : 134

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FIGURE 64 (cont)

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	*	560	*	580	*	600	
TrDFRh1 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAG-----						:586
TrDFRh2 :	AGAGTCAAAATGACCGGTTGGATGTATTTT-----						:569
TrDFRh3 :	-----						: -
TrDFRh4 :	ANAGTCAANATGACCGGTTGGATGTATTTTGTTCAAANACCC-----						:580
TrDFRh5 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCT-----						:581
TrDFRh6 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAG-----						:592
TrDFRh7 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT						:597
TrDFRh8 :	AGAGTCAAAATGACCGGTTGGATGTATTTTGTTCAAAGA-----						:576
TrDFRh9 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAG-----						:587
TrDFRh10 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT						:596
TrDFRh11 :	AGAGTCAAAATGACCGGTTGGATGTATTTTGTTCAAAGACCCT-----						:579
TrDFRh12 :	AGAGTCAAGATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT						:595
TrDFRh13 :	AGAGTCAAAATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAG-----						:586
TrDFRh14 :	AGAGTCAAAATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT						:552
TrDFRh15 :	AGAGTCAAAATGACCGGTTGGATGTATTTTGTTCAAAGACCCTAGCAGAGCAAGAAGCT						:194

	*	620	*	640	*	660	
TrDFRh1 :	-----						: -
TrDFRh2 :	-----						: -
TrDFRh3 :	-----						: -
TrDFRh4 :	-----						: -
TrDFRh5 :	-----						: -
TrDFRh6 :	-----						: -
TrDFRh7 :	TT-----						:598
TrDFRh8 :	-----						: -
TrDFRh9 :	-----						: -
TrDFRh10 :	TGGAAGTATTCCTAAAGAGCACAACATAGACTTTTG-----						:630
TrDFRh11 :	-----						: -
TrDFRh12 :	TGGAAGTAT-----						:604
TrDFRh13 :	-----						: -
TrDFRh14 :	TGGAAGTATTCGAAAGAGCACAACATAGACTTTTG-----						:586
TrDFRh15 :	TGGAAGTATTCGAAAGAGCACAACATAGACTTTGTCTCCATCATTCACCTCTTGTGTT						:254

	*	680	*	700	*	720	
TrDFRh1 :	-----						: -
TrDFRh2 :	-----						: -
TrDFRh3 :	-----						: -
TrDFRh4 :	-----						: -
TrDFRh5 :	-----						: -
TrDFRh6 :	-----						: -
TrDFRh7 :	-----						: -
TrDFRh8 :	-----						: -
TrDFRh9 :	-----						: -
TrDFRh10 :	-----						: -
TrDFRh11 :	-----						: -
TrDFRh12 :	-----						: -
TrDFRh13 :	-----						: -
TrDFRh14 :	-----						: -
TrDFRh15 :	GGCCCCTTTCTTATGGCCTCAATGCCACCTAGTCTAATCACTGCTCTTTCTCTTATCACA						:314

FIGURE 64 (cont)

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	*	740	*	760	*	780	
TrDFRh1 :	----		----		----		:
TrDFRh2 :	----		----		----		:
TrDFRh3 :	----		----		----		:
TrDFRh4 :	----		----		----		:
TrDFRh5 :	----		----		----		:
TrDFRh6 :	----		----		----		:
TrDFRh7 :	----		----		----		:
TrDFRh8 :	----		----		----		:
TrDFRh9 :	----		----		----		:
TrDFRh10 :	----		----		----		:
TrDFRh11 :	----		----		----		:
TrDFRh12 :	----		----		----		:
TrDFRh13 :	----		----		----		:
TrDFRh14 :	----		----		----		:
TrDFRh15 :	GGAAATGAGGCCCAT	T	TACTCAATCATAAAGCAAGGGCAATACGTCCATTTAGATGACCTT				: 374

	*	800	*	820	*	840	
TrDFRh1 :	----		----		----		:
TrDFRh2 :	----		----		----		:
TrDFRh3 :	----		----		----		:
TrDFRh4 :	----		----		----		:
TrDFRh5 :	----		----		----		:
TrDFRh6 :	----		----		----		:
TrDFRh7 :	----		----		----		:
TrDFRh8 :	----		----		----		:
TrDFRh9 :	----		----		----		:
TrDFRh10 :	----		----		----		:
TrDFRh11 :	----		----		----		:
TrDFRh12 :	----		----		----		:
TrDFRh13 :	----		----		----		:
TrDFRh14 :	----		----		----		:
TrDFRh15 :	TGTCTTGCTCATATATTTCTGTATGAGAATCCAAAAGCTCAAGGGAGATACATTTGCTGT						: 434

	*	860	*	880	*	900	
TrDFRh1 :	----		----		----		:
TrDFRh2 :	----		----		----		:
TrDFRh3 :	----		----		----		:
TrDFRh4 :	----		----		----		:
TrDFRh5 :	----		----		----		:
TrDFRh6 :	----		----		----		:
TrDFRh7 :	----		----		----		:
TrDFRh8 :	----		----		----		:
TrDFRh9 :	----		----		----		:
TrDFRh10 :	----		----		----		:
TrDFRh11 :	----		----		----		:
TrDFRh12 :	----		----		----		:
TrDFRh13 :	----		----		----		:
TrDFRh14 :	----		----		----		:
TrDFRh15 :	TCACATGAAGCAACCATT	CATCAAGTTGCAAAACTTATTAAAGAAAAATACCCAGAGTTC					: 494

FIGURE 64 (cont)

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	*	920	*	940	*	960	
TrDFRh1 :	-----		-----		-----		:
TrDFRh2 :	-----		-----		-----		:
TrDFRh3 :	-----		-----		-----		:
TrDFRh4 :	-----		-----		-----		:
TrDFRh5 :	-----		-----		-----		:
TrDFRh6 :	-----		-----		-----		:
TrDFRh7 :	-----		-----		-----		:
TrDFRh8 :	-----		-----		-----		:
TrDFRh9 :	-----		-----		-----		:
TrDFRh10 :	-----		-----		-----		:
TrDFRh11 :	-----		-----		-----		:
TrDFRh12 :	-----		-----		-----		:
TrDFRh13 :	-----		-----		-----		:
TrDFRh14 :	-----		-----		-----		:
TrDFRh15 :	<b>AATGTCCCAACAAAATTCAATGATATCCCAGATGAATTGGAAATTATTAAATTTTCTAAA</b>						: 554

	*	
TrDFRh1 :	-----	:
TrDFRh2 :	-----	:
TrDFRh3 :	-----	:
TrDFRh4 :	-----	:
TrDFRh5 :	-----	:
TrDFRh6 :	-----	:
TrDFRh7 :	-----	:
TrDFRh8 :	-----	:
TrDFRh9 :	-----	:
TrDFRh10 :	-----	:
TrDFRh11 :	-----	:
TrDFRh12 :	-----	:
TrDFRh13 :	-----	:
TrDFRh14 :	-----	:
TrDFRh15 :	<b>AAGAAGATCACAGACT</b>	: 570

## FIGURE 64 (cont)



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      *           20           *           40           *           60
TrLCRa : GGNCATAAAAAGTGCAGTGTGTATAAGTTTNTTAGTGAAAAAGAGTGTGTAAATTA : 60

      *           80           *           100          *           120
TrLCRa : ACATCATGGCTAGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG : 120

      *           140          *           160          *           180
TrLCRa : GTTTTGTTCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA : 180

      *           200          *           220          *           240
TrLCRa : CTGTTAGAGACCCAGATAGTCCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG : 240

      *           260          *           280          *           300
TrLCRa : GGGAAGTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG : 300

      *           320          *           340          *           360
TrLCRa : CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG : 360

      *           380          *           400          *           420
TrLCRa : AGAATGACATGATAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA : 420

      *           440          *           460          *           480
TrLCRa : GAGCAAAAGAAGTCAAAAGAGTTATCTTAACATCTTCGGCAGCCGCGGTGACTATAAATG : 480

      *           500          *           520          *           540
TrLCRa : AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA : 540

      *           560          *           580          *           600
TrLCRa : AACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAATGCTAGCTGAAAAGGCTG : 600

      *           620          *           640          *           660
TrLCRa : CATGGAATTTGCTGAAGAAAAATGACATTGATCTAATCACTGTGATACCTAGTTTAACAA : 660

      *           680          *           700          *           720
TrLCRa : CTGGTCCTTCTCTCACACCAGATATCCCATCTAGTGTTGGCTTGGCAATGTCTCTAATAA : 720

      *           740          *           760          *           780
TrLCRa : CAGGCAATGATTTCTCATAAATGCTCTGAAAGGAATGCAATTTCTGTGCGGTTCTGTTAT : 780

      *           800          *           820          *
TrLCRa : CCATCACTCATGTTGAGGATATTTGCCGAGCTCATATATTTCTGGCAGAGAAG : 833

```

## FIGURE 65

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TrLCRa :           \*          20          \*          40          \*          60  
MASIKQIGNKKACVIGGTGFVASMLIKQLLEKGYAVNTTVRDPDSPKKISHLVALQSLGE : 60

TrLCRa :           \*          80          \*          100          \*          120  
LNLFRADLTVEEDFDAPIAGCELVFQLATPVNFASQDPENDMIKPAIKGVLMNVLKASARA : 120

TrLCRa :           \*          140          \*          160          \*          180  
KEVKRVILTSSAAAVTINELKGTGHVMDETNWSDFEFLNTAKPPTWGYPASKMLAEKAAW : 180

TrLCRa :           \*          200          \*          220          \*          240  
KFAEENDIDLITVIPSLTTGPSLTPDIPSSVGLAMSLITGNDFLINALKGMQFLSGSLSI : 240

TrLCRa :           \*  
THVEDICRAHIFLAEK : 256

## FIGURE 66

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		*	20	*	40	*	60	
TrLCRa1:	GGNCATAAAAACTGCACTAGTGTGTATAAGTTTNTAGTGAAAAAAGAGTGTGTAAATTA	:	60					
TrLCRa2:	GGNCATAAAAACTGCACTAGTGTGTATAAGTTTNTAGTGAAAAAAGAGTGTGTAAATTA	:	60					
TrLCRa3:	-----TAAAAACTGTACTNGTGTGTATAAGTTTNTAGTGAAAAAAGAGTGTGTAAATTA	:	55					
TrLCRa4:	-----TAAAAACTGCACTAGTGTGTATAAGTTTCTTGCTGAAAAAAGAGTTTGTAAATTA	:	55					
TrLCRa5:	-----GAGCTCGTGTGNANTAGTTTCTTGCTGAAAAAAGAGTTTGTAAANTA	:	47					
TrLCRa6:	-----	:	-					
TrLCRa7:	-----	:	-					
		*	80	*	100	*	120	
TrLCRa1:	ACATCATGGCTAGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG	:	120					
TrLCRa2:	ACATCATGGCTAGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG	:	120					
TrLCRa3:	ACATCATGGCTAGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG	:	115					
TrLCRa4:	ACATCATGGCTAGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG	:	115					
TrLCRa5:	ACATCNTGGCTNGTATCAAACAAATTGGAAACAAGAAAGCATGTGTGATTGGTGGCACTG	:	107					
TrLCRa6:	-----	:	-					
TrLCRa7:	-----	:	-					
		*	140	*	160	*	180	
TrLCRa1:	GTTTTGTTGCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA	:	180					
TrLCRa2:	GTTTTGTTGCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA	:	180					
TrLCRa3:	GTTTTGTTGCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA	:	175					
TrLCRa4:	GTTTTGTTGCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA	:	175					
TrLCRa5:	GTTTTGTTGCATCTATGTTGATCAAACAGTTACTTGAAAAGGGTTATGCTGTTAATACTA	:	167					
TrLCRa6:	-----	:	-					
TrLCRa7:	-----	:	-					
		*	200	*	220	*	240	
TrLCRa1:	CTGTTAGAGACCCAGATAGTCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG	:	240					
TrLCRa2:	CTGTTAGAGACCCAGATAGTCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG	:	240					
TrLCRa3:	CTGTTAGAGACCCAGATAGTCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG	:	235					
TrLCRa4:	CCTGTTAGAGACCCAGATAGTCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG	:	235					
TrLCRa5:	CCTGTTAGAGACCCAGATAGTCCTAAGAAAATATCTCACCTAGTGGCACTGCAAAGTTTGG	:	227					
TrLCRa6:	-----G	:	1					
TrLCRa7:	-----	:	-					
		*	260	*	280	*	300	
TrLCRa1:	GGGAACCTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	300					
TrLCRa2:	GGGAACCTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	300					
TrLCRa3:	GGGAACCTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	295					
TrLCRa4:	GGGAACCTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	295					
TrLCRa5:	GGGAACCTGAATCTATTTAGAGCAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	287					
TrLCRa6:	NGNACCTGAATCTATTTAGAGNAGACTTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	61					
TrLCRa7:	-----TTAACAGTTGAAGAAGATTTTGATGCTCCTATAG	:	34					
		*	320	*	340	*	360	
TrLCRa1:	CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	360					
TrLCRa2:	CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	360					
TrLCRa3:	CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	355					
TrLCRa4:	CAGGATGTGAACTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	355					
TrLCRa5:	CAGGATGTGAACTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	347					
TrLCRa6:	CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	121					
TrLCRa7:	CAGGATGTGAGCTTGTTTTTCAACTTGCTACACCTGTGAACTTTGCTTCTCAAGATCCTG	:	94					

FIGURE 67

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	*	380	*	400	*	420	
TrLCRa1:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA						: 420
TrLCRa2:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA						: 420
TrLCRa3:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA						: 415
TrLCRa4:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAATTCGCAA						: 415
TrLCRa5:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAATTCGCAA						: 407
TrLCRa6:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA						: 181
TrLCRa7:	AGAATGACATGATAAAAGCCAGCAATCAAAGGTGTGTTGAATGTGTTGAAAGCAAGTGCAA						: 154

	*	440	*	460	*	480	
TrLCRa1:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 480
TrLCRa2:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 480
TrLCRa3:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 475
TrLCRa4:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 475
TrLCRa5:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 467
TrLCRa6:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 241
TrLCRa7:	GAGCAAAAGAAGTCAAAAGAGTTATCTTAAACATCTTCGGCAGCCGCGGTGACTATAAAATG						: 214

	*	500	*	520	*	540	
TrLCRa1:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 540
TrLCRa2:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 540
TrLCRa3:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 535
TrLCRa4:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 535
TrLCRa5:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 527
TrLCRa6:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 301
TrLCRa7:	AACTCAAAGGGACAGGTCATGTTATGGATGAAACCAACTGGTCAGATGTTGAATTTCTGA						: 274

	*	560	*	580	*	600	
TrLCRa1:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTT-----						: 576
TrLCRa2:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAN-----						: 580
TrLCRa3:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAAT-----						: 578
TrLCRa4:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAATGCTAGCTGAAAAGGCTG						: 595
TrLCRa5:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAATGCTAGCTGAAAAGGCTG						: 587
TrLCRa6:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAATGCTAGCTGAAAAGGCTG						: 361
TrLCRa7:	ACACTGCAAAGCCACCCACTTGGGGTTATCCTGCTTCAAAAATGCTAGCTGAAAAGGCTG						: 334

	*	620	*	640	*	660	
TrLCRa1:	-----						: -
TrLCRa2:	-----						: -
TrLCRa3:	-----						: -
TrLCRa4:	CATGGAAATTTGCTGAAGAAAATC-----						: 619
TrLCRa5:	CATGGAAATTTGCTGAAGAAAATGACNTTGAT-----						: 619
TrLCRa6:	CATGGAAATTTGCTGAAGAAAATGACATTGATCTAATCACTGTGATACCTAGTTTAACAA						: 421
TrLCRa7:	CATGGAAATTTGCTGAAGAAAATGACATTGATCTAATCACTGTGATACCTAGTTTAACAA						: 394

	*	680	*	700	*	720	
TrLCRa1:	-----						: -
TrLCRa2:	-----						: -
TrLCRa3:	-----						: -
TrLCRa4:	-----						: -
TrLCRa5:	-----						: -
TrLCRa6:	CTGGTCCTTCTCTCACACCAGATATCCCATCTAGTGTGGCTTGGCAATGTCTCTAATAA						: 481
TrLCRa7:	CTGGTCCTTCTCTCACACCAGATATCCCATCTAGTGTGGCTTGGCAATGTCTCTAATAA						: 454

FIGURE 67 (cont)

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	*	740	*	760	*	780		
TrLCRa1:	-----		-----		-----		:	-
TrLCRa2:	-----		-----		-----		:	-
TrLCRa3:	-----		-----		-----		:	-
TrLCRa4:	-----		-----		-----		:	-
TrLCRa5:	-----		-----		-----		:	-
TrLCRa6:	CAGGCAATGATTTCCCTCATAAATGCTCTGAAAGGAATGCAATTTCTGTCGGGTTTCGTTAT						:	541
TrLCRa7:	CAGGCAATGATTTCCCTCATAAATGCTCTGAAAGGAATGCAATTTCTGTCGGGTTTCGTTAT						:	514

	*	800	*	820	*		
TrLCRa1:	-----		-----		-----	:	-
TrLCRa2:	-----		-----		-----	:	-
TrLCRa3:	-----		-----		-----	:	-
TrLCRa4:	-----		-----		-----	:	-
TrLCRa5:	-----		-----		-----	:	-
TrLCRa6:	CCATCACTCATGTTGAGGATATTTGCCGAGCTCATATATTTCTGC					:	586
TrLCRa7:	CCATCACTCATGTTGAGGATATTTGCCGAGCTCATATATTTCTGGCAGAGAAG					:	567

**FIGURE 67 (cont)**

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TrF3'5'Ha: GGAACCAATTTGTCTCGGACTTTTTCCTCCGGGTTGGCCCGATTTCGATTTGCAGGGTGTGGTG : 60

TrF3'5'Ha: AAAGAGATGGATGTCTTGGTTCCACGTTTTGATAGCATATTTGAAAAAATGATTGGTGAA : 120

TrF3'5'Ha: CGTAAGAAGAAGGAAGTGGAGGGGAAAGAAAATGAAAGTAAGGATTTTCTGCAGTTTTTG : 180

TrF3'5'Ha: TTGAATTTGAAGGATGAGGGTGATTCTAAGACTCCATTCACAATTACCCATGTTAAGGCT : 240

TrF3'5'Ha: CTAATCATGGACATGGTTTGTGGGTGGATCAGACACATCCTCCAACACAATTGAGTTTGCA : 300

TrF3'5'Ha: TTGGCAGAAATGATGAACAACCCAGAAGTAATGAGGAAGGTTCAAGAGGAATTAGAAGAT : 360

TrF3'5'Ha: GTAGTTGGGAAAGATAACTTAGTAGAAGAGTCTCACATTCATAAGCTACCCCTACTTGCAT : 420

TrF3'5'Ha: GCAGTGATGAAAGAAACACTTCGTTTACACCCAGCACTTCCACTTTTAGTCCCTCACTGT : 480

TrF3'5'Ha: CCAAGTGAAACCACCAATGTTGGAGGCTACACAATTCCAAAGGGATCTCGTGTGTTTGTG : 540

TrF3'5'Ha: AACGTTTGGGCTATTCATAGAGACCCTTCATTTGGGAGAAACCACTAGAATTTGAT : 597

## FIGURE 68

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```

      *           20           *           40           *           60
TrF3'5'Ha: GTNLSDFFPGLARFDLQGVVKEMDVLVPRFDSIFEKMI GERKKKEVEGKENESKDFLQFL : 60

      *           80           *           100          *           120
TrF3'5'Ha: LNLKDEGDSKTPFTITHVKALLMDMVVGGSDTSSNTIEFALAEMMNNPEVMRKVQEELED :120

      *           140          *           160          *           180
TrF3'5'Ha: VVGKDNLVEESHIHKLPYLHAVMKETLRLHPALPLLVP HCPSETTNVGGYTIPKGSRVFV :180

      *
TrF3'5'Ha: NVWAIHRDPSIWEKPLEFD :199
```

## FIGURE 69

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TrF3 '5' Hb: GNAATCCACNAATCTCTTGAANTAATACCATTCTTTACAAGAACTTAACCATGGTGATG : 60

TrF3 '5' Hb: ATCACTCAATACCAAACCTTCCTTTACAAAGAACTTTCTATATCCTTTTTCATTTTCTTG : 120

TrF3 '5' Hb: ATAACCCATTTTCATCATAAGTTTCTCTTCAAAAAAATCTCAAAAAACTTCCACCAGGC : 180

TrF3 '5' Hb: CCAAAGGTTTTCAGTTGTTGGTGCACCTCCACTAATGGGATCCATGCCTCATGTTACC : 240

TrF3 '5' Hb: CTATTCAAAATGTCACAAAAATATGGTCCCATAATGTACCTAAAAATGGGATCAAATAAC : 300

TrF3 '5' Hb: ATGGTTGTAGCATCAACTCCTTCTTCAGCCAAAGCATTCTCTCAAAACACTTGACCTAAAT : 360

TrF3 '5' Hb: TTCTCCAATAGACCGCCGAACGCTGGCGCAACTCACCTAGCTTATGATTCACAAGACTTG : 420

TrF3 '5' Hb: GTTTTCGCCGACTATGGATCTAGGTGGAAATTACTTAGGAACTAAGTAAGTGCACATG : 480

TrF3 '5' Hb: CTCGGCGGAAAAGCCCTCGAAAATTGGTTCGAAAGTTCGTGAGATTGAAATGGGTCACATG : 540

TrF3 '5' Hb: ATTCGTACAATGTACGATTGTAGCAAGAAAGACGAATCCGTTGTTGTGGCCGAAATGTTG : 600

TrF3 '5' Hb: ACATATGCTATGGCCAATATGATAGGTCAAGTTATATTGAGTCGTCGCGTGTTCGAGACA : 660

TrF3 '5' Hb: AAAGGTAGTGACTCAAATGAATTTAAGGATATGGTTGNTG : 700

## FIGURE 70



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```

      *      20      *      40      *      60
TrF3'5'Hb: MVMITQYQTFLYKELSSISFFIFLITHFIISFLFKKNLKKLPPGPKGFPVVGALPLMGSMF : 60

      *      80      *      100     *      120
TrF3'5'Hb: HVTLFKMSQKYGPIMYLMGSSNNMVVASTPSSAKAFLKTLDLNFSSNRPPNAGATHLAYDS :120

      *      140     *      160     *      180
TrF3'5'Hb: QDLVFADYGSRWKLLRKLSNLHMLGGKALENWSKVREIEMGHMIRTMYDCSKKDESVVVA :180

      *      200     *
TrF3'5'Hb: EMLTYAMANMIGQVILSRRVFETKGSDSNEFKDMVX :216

```

## FIGURE 71

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```

      *           20           *           40           *           60
TrF3 '5' Hb1: GNAATCCACNAATCTCTTGAAATAATNCCATTTCCTTTACAAGAACTTAACCATGGTGATG : 60
TrF3 '5' Hb2: -----GNCNAATCTCTTGCANTAAANNCCATTTCCTTTACAAGAACTTAACCATGGTGATG : 54
TrF3 '5' Hb3: -----CNAATCTCTTGAANTNATACCATTTCCTTTACAAGAACTTAACCNITGGTGATG : 52
TrF3 '5' Hb4: -----TCTCTTCNAATNATACCATTTCCTTTACAAGAACTTAACCNITGGTGATG : 48

      *           80           *           100          *           120
TrF3 '5' Hb1: ATCACTCAATACCAAACCTTCCTTTCAAAGAACTTTCATATTCCTTTTTTCATTTTCTTG : 120
TrF3 '5' Hb2: ATCACTCAAACCAAACCTTCCTTTCAAAGAACTTTCATATTCCTTTTTTCATTTTCTTG : 114
TrF3 '5' Hb3: ATNNCTCAATACCAAACCTTCCTTTCAAAGAACTTTCATATTCCTTTTTTCATTTTCTTG : 112
TrF3 '5' Hb4: ATNNCTCNATACCAAACCTTCCTTTCAAAGAACTTTCATATTCCTTTTTTCATTTTCTTG : 108

      *           140          *           160          *           180
TrF3 '5' Hb1: ATAACCCATTTTCATCATAAGTTTCTCTTCAAAAAAAATCTCAAAAAACTTCCACCAGGC : 180
TrF3 '5' Hb2: ATAACCCATTTTCATCATAGTTTCTCTTCAAAAAAAATCTCAAAAAACTTCCACCAGGC : 174
TrF3 '5' Hb3: ATAACCCATTTTCATCATAAGTTTCTCTTCAAAAAAAATCTCAAAAAACTTCCACCAGGC : 172
TrF3 '5' Hb4: ATAACCCATTTTCATCATAGTTTCTCTTCAAAAAAAATCTCAAAAAACTTCCACCAGGC : 168

      *           200          *           220          *           240
TrF3 '5' Hb1: CCAAAAGGTTTTCCAGTTGTTGGTGCACCTCCACTAATGGGATCCATGCCTCATGTTACC : 240
TrF3 '5' Hb2: CCAAAAGGTTTTCCAGTTGTTGGTGCACCTCCACTAATGGGATCCATGCCTCATGTTACC : 234
TrF3 '5' Hb3: CCAAAAGGTTTTCCAGTTGTTGGTGCACCTCCACTAATGGGATCCATGCCTCATGTTACC : 232
TrF3 '5' Hb4: CCAAAAGGTTTTCCAGTTGTTGGTGCACCTCCACTAATGGGATCCATGCCTCATGTTACC : 228

      *           260          *           280          *           300
TrF3 '5' Hb1: CTATTCAAAATGTCACAAAAATATGGTCCCATAATGTACCTAAAAATGGGATCAAATAAGC : 300
TrF3 '5' Hb2: CTATTCAAAATGTCACAAAAATATGGTCCCATAATGTACCTAAAAATGGGATCAAATAAC : 294
TrF3 '5' Hb3: CTATTCAAAATGTCACAAAAATATGGTCCCATAATGTACCTAAAAATGGGATCAAATAAC : 292
TrF3 '5' Hb4: CTATTCAAAATGTCACAAAAATATGGTCCCATAATGTACCTAAAAATGGGATCAAATAAC : 288

      *           320          *           340          *           360
TrF3 '5' Hb1: ATGGTTGTAGCATCAACTCCTTCTTCAGCCAAAGCATTTCTCAAAACACTTGACCTAAAT : 360
TrF3 '5' Hb2: ATGGTTGTAGCATCAACTCCTTCTTCAGCCAAAGCATTTCTCAAAACACTTGACCTAAAT : 354
TrF3 '5' Hb3: ATGGTTGTAGCATCAACTCCTTCTTCAGCCAAAGCATTTCTCAAAACACTTGACCTAAAT : 352
TrF3 '5' Hb4: ATGGTTGTAGCATCAACTCCTTCTTCAGCCAAAGCATTTCTCAAAACACTTGACCTAAAT : 348

      *           380          *           400          *           420
TrF3 '5' Hb1: TTCTCCAATAGACCGCCGAACGCTGGCGCACTCACCTAGCTTATGATTCACAAGACTTG : 420
TrF3 '5' Hb2: TTCTCCAATAGACCGCCGAACGCTGGCGCACTCACCTAGCTTATGATTCACAAGACTTG : 414
TrF3 '5' Hb3: TTCTCCAATAGACCGCCGAACGCTGGCGCACTCACCTAGCTTATGATTCACAAGACTTG : 412
TrF3 '5' Hb4: TTCTCCAATAGACCGCCGAACGCTGGCGCACTCACCTAGCTTATGATTCACAAGACTTG : 408

      *           440          *           460          *           480
TrF3 '5' Hb1: GTTTTCGCCGACTATGGATCTAGGTGGAATTACTTAGGAAACTAAGTAACCTTGCACATG : 480
TrF3 '5' Hb2: GTTTTCGCCGACTATGGATCTAGGTGGAATTACTTAGGAAACTAAGTAACCTTGCACATG : 474
TrF3 '5' Hb3: GTTTTCGCCGACTATGGATCTAGGTGGAATTACTTAGGAAACTAAGTAACCTTGCACATG : 472
TrF3 '5' Hb4: GTTTTCGCCGACTATGGATCTAGGTGGAATTACTTAGGAAACTAAGTAACCTTGCACATG : 468

      *           500          *           520          *           540
TrF3 '5' Hb1: CTCGGCGGAAAAGCCCTCGAAATTTGGTTCGAAAGTTCGTGAGATTGAAATGGGTACATG : 540
TrF3 '5' Hb2: CTCGGCGGAAAAGCCCTCGAAATTTGGTTCGAAAGTTCGTGAGATTGAAATGGGTACATG : 534
TrF3 '5' Hb3: CTCGGCGGAAAAGCCCTCGAAATTTGGTTCGAAAGTTCGTGAGATTGAAATGGGTACATG : 532
TrF3 '5' Hb4: CTCGGCGGAAAAGCCCTCGAAATTTGGTTCGAAAGTTCGTGAGATTGAAATGGGTACATG : 528

```

FIGURE 72

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```

      *           560           *           580           *           600
TrF3'5'Hb1: ATTCGTACAATGTATGATTGTAGCAAGAAAGACGAATCCGTTGT----- : 584
TrF3'5'Hb2: ATTCGTACAATGTACGATTGTAGCAAGAAAGACGAATCCGTTGTTGTGGCCGAAATGTTG : 594
TrF3'5'Hb3: ATTCGTACAATGTACGATTGTAGCAAGAAAGACGAATATGTTGTTGTC----- : 580
TrF3'5'Hb4: ATTCGTACAATGTACGATTGTAGCAAGAAAGACGAATCCGNTGNTG----- : 74

      *           620           *           640           *           660
TrF3'5'Hb1: ----- : -
TrF3'5'Hb2: ACATATGCTATGGCCAATATGATAGGTCAAGTTATATTGAGTCGTCGCGTGTTCGAGACA : 654
TrF3'5'Hb3: ----- : -
TrF3'5'Hb4: ----- : -

      *           680           *           700
TrF3'5'Hb1: ----- : -
TrF3'5'Hb2: AAAGGTAGTGA CTCAAATGAATTTAAGGATATGGTTGNTG : 694
TrF3'5'Hb3: ----- : -
TrF3'5'Hb4: ----- : -
```

## FIGURE 72 (cont)

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```

      *           20           *           40           *           60
TrF3Ha : GCACACNTCTATTTATTTCTACTTAAACCTNACAAAAATAANACCCACAAAACACAAAC :   60

      *           80           *           100          *           120
TrF3Ha : ACCACAAACACCAAAACCGAGTCCGTTTCCTNNTCNAACATGGCACCAAGCCAAACTCTA :  120

      *           140          *           160          *           180
TrF3Ha : AGTTATCTCTCACAACAAAACACTCTTGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT :  180

      *           200          *           220          *           240
TrF3Ha : CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCTCTTGCTGGAATT :  240

      *           260          *           280          *           300
TrF3Ha : GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT :  300

      *           320          *           340          *           360
TrF3Ha : TGGGGTATTTTTCAGGTTGTTGATCATGGTGTTGATACAAAACCTGTTTCTGAGATGACC :  360

      *           380          *           400          *           420
TrF3Ha : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :  420

      *           440          *           460          *           480
TrF3Ha : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :  480

      *           500          *           520          *           540
TrF3Ha : TGGAGAGAGCTAGTGACATATTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG :  540

      *           560          *           580          *           600
TrF3Ha : CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT :  600

      *           620          *           640          *           660
TrF3Ha : TTAGCTTGCAAACCTATTGGAAGTTTTATCAGAAGCAATGGGGTTTAGAAAAAGAAGCTCTA :  660

      *           680          *           700          *           720
TrF3Ha : ACAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT :  720

      *           740          *           760          *           780
TrF3Ha : GAACCTGACCTCACACTTGGCCTTAAACGTCACACTGACCCTGGCACAATTACTCTTTTG :  780

```

## FIGURE 73

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```

      *           800           *           820           *           840
TrF3Ha : CTTCAAGATCAAGTTGGTGGCCTTCAAGCTACCAAAGATAATGGTAAGACGTGGATTACA : 840

      *           860           *           880           *           900
TrF3Ha : GTTCAACCAGTTGAAGGTGCTTTTGTGTGTTAATCTTGGAGACCATGGTCACTATCTAAGT : 900

      *           920           *           940           *           960
TrF3Ha : AATGGACGGTTCAAAAATGCTGACCACCAAGCAGTGGTGAATTCGAACTACAGCCGNTTA : 960

      *           980           *          1000           *          1020
TrF3Ha : TCAATAGCAACATTTCAAAATCCAGCTCCCGATGCAACTGTATACCCTTTGAAGATTAGA : 1020

      *          1040           *          1060           *          1080
TrF3Ha : GAGGGTGAAAAATCTGTGTTGGAAGAACCAATCACTTTTGCTGAAATGTATAGAAGGAAG : 1080

      *          1100           *          1120           *          1140
TrF3Ha : ATGACCAAAGACCTTGAAATTGCTAGGATGAAGAAGTTGGCTAAGGAACAACAACCTTAGG : 1140

      *          1160           *          1180           *          1200
TrF3Ha : GACTTGGAGGAGAACAAGACTAAATATGAGGCCAAACCTTTGAATGAGATCTTTGCTTAA : 1200

      *          1220           *          1240           *          1260
TrF3Ha : TTAATTAGTCTTAATTTAAATAATTAATAAATTTAGACTTAATTTACATATAATAATTT : 1260

TrF3Ha : T : 1261

```

## FIGURE 73 (cont)

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```

      *      20      *      40      *      6
TrF3Ha : MAPSQTL SYLSQQNTLESSFVREEDERPKVAYNNFSNEIPIISLAGIDEVDGRRTEICNK : 59

      *      80      *      100     *      120
TrF3Ha : IVEACENWGIFQVVDHGVDTKLVSEMTRFAREFFALPPEEKLRFDMSGGKKGGFIVSSHL : 120

      *      140     *      160     *      180
TrF3Ha : QGEAVKDWRELVTYFSYPIKQRDYSRWPDKPEGWKEVTEKYSENLMNLACKLLEVLSEAM : 180

      *      200     *      220     *      240
TrF3Ha : GLEKEALTKACVDMDQKVVINYYPKCPEPDLTLGLKRHTDPGTITLLQLDQVGGLQATKD : 240

      *      260     *      280     *      300
TrF3Ha : NGKTWITVQPVGEAFVVNLGDHGHYLSNGRFGKNADHQAVVNSNYSXLSIATFQNPAPDAT : 300

      *      320     *      340     *      360
TrF3Ha : VYPLKIREGEKSVLEEPITFAEMYRRKMTKDLEIARMKKLAKEQQLRDLEENKTKYEAKP : 360

TrF3Ha : LNEIFA : 366

```

## FIGURE 74

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		*	20	*	40	*	60	
TrF3Ha1	:	GCACACNTCTATTTATTTTCTACTTAAACCT	CACAAAAAATAA	-	ACCCACAACACACAAC	:	59	
TrF3Ha2	:	GCACACNTCTATTTATTTTCTACTTAAACCT	---NAAAAAATAA	-	ACCCACAA--CAC-AA-	:	52	
TrF3Ha3	:	-CACACNTCTATTTATTTTCTACTTAAACCT	NACAAAAAATAAN	ACCCACAACACACAAC	:	59		
TrF3Ha4	:	--TCCCTCTANTTATTTCTNCTTAAACCT	NNCAAAAAATNANN	ACCCACAACACACNAAN	:	58		
TrF3Ha5	:	-----	-----	-----	GTTAACACAC-N-CAACACAAC	:	22	
TrF3Ha6	:	-----	-----	-----	CNA-ACCCACAACACACAAC	:	20	
TrF3Ha7	:	-----	-----	-----	CCACNAAN-N-CAAAC	:	16	
TrF3Ha8	:	-----	-----	-----	CACNATNACACAAC	:	16	
TrF3Ha9	:	-----	-----	-----	CCACANGACAC-AA-	:	13	
TrF3Ha10	:	-----	-----	-----	GAANGCCACNAAC	:	13	
TrF3Ha11	:	-----	-----	-----	GNACGACACAAC	:	13	
TrF3Ha12	:	-----	-----	-----	GGGA--CAC-AA-	:	9	
TrF3Ha13	:	-----	-----	-----	T-TAACACAAC	:	12	
TrF3Ha14	:	-----	-----	-----	CNCAACACAAC	:	12	
TrF3Ha15	:	-----	-----	-----	GNAATC-AA-	:	11	
TrF3Ha16	:	-----	-----	-----	GAAC-C-AA-	:	10	
TrF3Ha17	:	-----	-----	-----	AACSCAAG	:	9	
TrF3Ha18	:	-----	-----	-----	GCACNAAC	:	8	
TrF3Ha19	:	-----	-----	-----	AC-ANC	:	5	
TrF3Ha20	:	-----	-----	-----	GCNNCA	:	6	
TrF3Ha21	:	-----	-----	-----	GNAC-C	:	6	
TrF3Ha22	:	-----	-----	-----	C-AA-AA	:	5	
TrF3Ha23	:	-----	-----	-----	AA-AA	:	4	
TrF3Ha24	:	-----	-----	-----	AA-AA	:	3	
TrF3Ha25	:	-----	-----	-----	AA-AA	:	3	
TrF3Ha26	:	-----	-----	-----	GC	:	2	
TrF3Ha27	:	-----	-----	-----	GC	:	2	
TrF3Ha28	:	-----	-----	-----	GC	:	2	
TrF3Ha29	:	-----	-----	-----	A	:	1	
TrF3Ha30	:	-----	-----	-----		:	-	
TrF3Ha31	:	-----	-----	-----		:	-	
TrF3Ha32	:	-----	-----	-----		:	-	
TrF3Ha33	:	-----	-----	-----		:	-	
TrF3Ha34	:	-----	-----	-----		:	-	
TrF3Ha35	:	-----	-----	-----		:	-	
TrF3Ha36	:	-----	-----	-----		:	-	
TrF3Ha37	:	-----	-----	-----		:	-	
TrF3Ha38	:	-----	-----	-----		:	-	
TrF3Ha39	:	-----	-----	-----		:	-	
TrF3Ha40	:	-----	-----	-----		:	-	
TrF3Ha41	:	-----	-----	-----		:	-	

FIGURE 75

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```

      *           80           *           100           *           120
TrF3Ha1 : ACCA AAAACAG-AG ACCGTTTCC TCAATC---A---AACATGGCACCAAGCCAAACTCTA :113
TrF3Ha2 : ACCA AAAACAG-AG ACCGTTTCCATCATC---A---AACATGGCACCAAGCCAAACTCTA :106
TrF3Ha3 : ACCA AAAACAGNAG ACCGTTTCCATCATC NNNNANNAACATGGCACCAAGCCAAACTCTA :119
TrF3Ha4 : ANCAC AAAANANACAGNAGTACCGTTTCCATCATCAACATGGCACCAAGCCAAACTCTA :118
TrF3Ha5 : ACCA AAAACAGNAG ACCGTTTCC-A-TNCA-T:CAAACATGGCACCAAGCCAAACTCTA : 78
TrF3Ha6 : A-CA AAAACAG-AG-NCCGTTTCN-TNATC---A---AACATGGCACCAAGCCAAACTCTA : 71
TrF3Ha7 : ACCA AAAACAGNAG ACCGTTTCC-A-TNCA-TNCAACATGGCACCAAGCCAAACTCTA : 73
TrF3Ha8 : ACCA AAAACAGNAG ACCGTTTCC-A-TNCA-T:CAAACATGGCACCAAGCCAAACTCTA : 72
TrF3Ha9 : A-CACCAAA-A-ACAGTAGCAACCGTTTCCATCATCAACATGGCACCAAGCCAAACTCTA : 70
TrF3Ha10 : ANCAAGACNCAAAAANAGAGACCGTTTCCCTNNNTNNAACATGGCACCAAGCCAAACTCTA : 73
TrF3Ha11 : ACCA AAAACAG-AG ACCGTTTCC-TNATC---A---AACATGGCACCAAGCCAAACTCTA : 66
TrF3Ha12 : ACC-AAAAC-G-AGC-CCGTTTCC-T-NATC---N---AACATGGCACCAAGCCAAACTCTA : 57
TrF3Ha13 : ACCA AAAACAG-AGNACCGTTTCC-TNTTC---A---A-CATGGCACCAAGCCAAACTCTA : 64
TrF3Ha14 : ACCA AAAACAG-AG ACCGTTTCC-A-TNCA-TNCAACATGGCACCAAGCCAAACTCTA : 69
TrF3Ha15 : AC-TNCAAAACCCA-AA-CGAG-CCGTTTCCCTNCTNNAACATGGCACCAAGCCAAACTCTA : 71
TrF3Ha16 : AC-CCAAACCCA-AA-CGAGNCCGTTTCCCT-NTC-AACATGGCACCAAGCCAAACTCTA : 67
TrF3Ha17 : AC-CAAAACCCA-AA-CGAG-CCGTTTCCCTNNNTNNAACATGGCACCAAGCCAAACTCTA : 69
TrF3Ha18 : AC-CACAAAC-CC-AAA-NGAGACCGTTTCCCT-NTNNAACATGGCACCAAGCCAAACTCTA : 68
TrF3Ha19 : A-CNNAAAC-N-C-AAA-CGAGTCCGTTTCCCT-NTNNAACATGGCACCAAGCCAAACTCTA : 60
TrF3Ha20 : AN-CACAAAC-C-AAAACGAGCAACCGTTTCCCTANTNNAACATGGCACCAAGCCAAACTCTA : 66
TrF3Ha21 : AC-TNCAAAACCCA-AA-CGAG-CCGTTTCCCTNCTNNAACATGGCACCAGCCAAACTCTA : 66
TrF3Ha22 : AC-TNNAAAACCCA-AA-CGAGNCCGTTTCCCT-CTC-AACATGGCACCAAGCCAAACTCTA : 62
TrF3Ha23 : AC-CAAAACCCA-AA-CGAG-CCGTTTCCCT-CTN-AACATGGCACCAAGCCAAACTCTA : 60
TrF3Ha24 : AC-CAAAACCCA-AA-CGAG-CCGTTTCCCT-NTC-AACATGGCACCAAGCCAAACTCTA : 58
TrF3Ha25 : AC-CAAAACCCA-AA-CGAGNCCGTTTCCCT-NTC-AACATGGCACCAAGCCAAACTCTA : 59
TrF3Ha26 : AC-CACAAAC-ACCAAAA-CGAG-CCGTTTCC-NTCAACATGGCACCAAGCCAAACTCTA : 59
TrF3Ha27 : AC-TNCAAAACCCA-AA-CGAG-CCGTTTCCCTNCTNNAACATGGCACCAAGCCAAACTCTA : 62
TrF3Ha28 : AC-TNCAAAACCCA-AA-CGAGNCCGTTTCCCTNNNTNNAACATGGCACCAAGCCAAACTCTA : 62
TrF3Ha29 : ACC-AAAACAG-AGNACCGTTTTCN-TNNNN---A---A-CATGGC-CC-AGCCAAACTCTA : 51
TrF3Ha30 : A-CAAAAC-C-T-NAAACCGAGACCGTTTCCCT-NTNNAACATGGCACCAAGCCAAACTCTA : 55
TrF3Ha31 : -GCNNAAACACCA-AA-CGAG-CCGTTTCCCTNNNTGAAACATGGCACCAAGCCAAACTCTA : 59
TrF3Ha32 : -CAACAAACCCA-AA-CGAG-CCGTTTCCCT-NTC-AACATGGCACCAAGCCAAACTCTA : 56
TrF3Ha33 : -----CAAAC-C-AAAACNGAGNCCGTTTCCCTA-TN-AACATGGCACCAAGCCAAACTCTA : 52
TrF3Ha34 : -----NATC-C-AAAACG-AGNCCGTTTCCCTNCTC-AACATGGCTCCTAGCCAAACTCTA : 51
TrF3Ha35 : -----GAAACCAAAACNGAG-CCGTTTTCNNNTNNAACATGGCACCAGCCAAACTCTA : 54
TrF3Ha36 : -----A-CACAAAC-AGAGNCCGTTTTCNNNTNNAACATGGCACCAGCCAAACTCTA : 53
TrF3Ha37 : -----AACCAAAACCGAGNCCGTTTCCNNNTNNAACATGGCACCAGCCAAACTCTA : 52
TrF3Ha38 : -----AACCAAAACCGAGNCCGTTTTCNNNTNNAACATGGCACCAGCCAAACTCTA : 52
TrF3Ha39 : ----- : 1
TrF3Ha40 : ----- : -
TrF3Ha41 : ----- : -

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FIGURE 75 (cont)



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	*	140	*	160	*	180	
TrF3Ha1	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	173			
TrF3Ha2	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	166			
TrF3Ha3	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	179			
TrF3Ha4	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	178			
TrF3Ha5	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	138			
TrF3Ha6	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	131			
TrF3Ha7	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	133			
TrF3Ha8	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	132			
TrF3Ha9	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	130			
TrF3Ha10	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	133			
TrF3Ha11	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	126			
TrF3Ha12	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	116			
TrF3Ha13	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	124			
TrF3Ha14	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	129			
TrF3Ha15	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	131			
TrF3Ha16	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	127			
TrF3Ha17	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	129			
TrF3Ha18	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	128			
TrF3Ha19	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	120			
TrF3Ha20	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	126			
TrF3Ha21	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	126			
TrF3Ha22	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	122			
TrF3Ha23	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	119			
TrF3Ha24	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	117			
TrF3Ha25	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	118			
TrF3Ha26	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	119			
TrF3Ha27	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	122			
TrF3Ha28	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	122			
TrF3Ha29	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	110			
TrF3Ha30	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	115			
TrF3Ha31	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	119			
TrF3Ha32	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	116			
TrF3Ha33	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	112			
TrF3Ha34	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	109			
TrF3Ha35	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	114			
TrF3Ha36	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	113			
TrF3Ha37	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	112			
TrF3Ha38	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	112			
TrF3Ha39	:	AGTTATCTCTCACAACAAAACACTCTCGAGTCAAGTTTCGTTAGGGAAGAAGATGAGCGT	:	61			
TrF3Ha40	:	-----	:	-			
TrF3Ha41	:	-----	:	-			

FIGURE 75 (cont)

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	*	200	*	220	*	240	
TrF3Ha1	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	233			
TrF3Ha2	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	226			
TrF3Ha3	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	239			
TrF3Ha4	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	238			
TrF3Ha5	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	198			
TrF3Ha6	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	191			
TrF3Ha7	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	193			
TrF3Ha8	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	192			
TrF3Ha9	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	190			
TrF3Ha10	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	193			
TrF3Ha11	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	186			
TrF3Ha12	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	176			
TrF3Ha13	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	184			
TrF3Ha14	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	189			
TrF3Ha15	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	191			
TrF3Ha16	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	187			
TrF3Ha17	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	189			
TrF3Ha18	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	188			
TrF3Ha19	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	180			
TrF3Ha20	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	186			
TrF3Ha21	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	186			
TrF3Ha22	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	182			
TrF3Ha23	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	179			
TrF3Ha24	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	177			
TrF3Ha25	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	178			
TrF3Ha26	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	179			
TrF3Ha27	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	182			
TrF3Ha28	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	182			
TrF3Ha29	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	170			
TrF3Ha30	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	175			
TrF3Ha31	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	179			
TrF3Ha32	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	176			
TrF3Ha33	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	172			
TrF3Ha34	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	169			
TrF3Ha35	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	174			
TrF3Ha36	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	173			
TrF3Ha37	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	172			
TrF3Ha38	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	172			
TrF3Ha39	:	CCAAAAGTTGCCTACAATAACTTCAGCAACGAGATTCCAATCATTTCCTCTTGCTGGAATT	:	121			
TrF3Ha40	:	-----	:	-			
TrF3Ha41	:	-----	:	-			

FIGURE 75 (cont)

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	*	260	*	280	*	300	
TrF3Ha1	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	293			
TrF3Ha2	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	286			
TrF3Ha3	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	299			
TrF3Ha4	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	298			
TrF3Ha5	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	258			
TrF3Ha6	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	251			
TrF3Ha7	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	253			
TrF3Ha8	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	252			
TrF3Ha9	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	250			
TrF3Ha10	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	253			
TrF3Ha11	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	246			
TrF3Ha12	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	236			
TrF3Ha13	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	244			
TrF3Ha14	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	249			
TrF3Ha15	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	251			
TrF3Ha16	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	247			
TrF3Ha17	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	249			
TrF3Ha18	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	236			
TrF3Ha19	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	240			
TrF3Ha20	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	246			
TrF3Ha21	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	246			
TrF3Ha22	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	242			
TrF3Ha23	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	239			
TrF3Ha24	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	237			
TrF3Ha25	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	238			
TrF3Ha26	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	239			
TrF3Ha27	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	242			
TrF3Ha28	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	242			
TrF3Ha29	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	230			
TrF3Ha30	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	235			
TrF3Ha31	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	239			
TrF3Ha32	:	GATGAGGTTGATGGTCGTAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	236			
TrF3Ha33	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	232			
TrF3Ha34	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	229			
TrF3Ha35	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	234			
TrF3Ha36	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	233			
TrF3Ha37	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	232			
TrF3Ha38	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	232			
TrF3Ha39	:	GATGAGGTTGATGGTCGGAAGAACAGAGATATGTAACAAGATTGTTGAAGCTTGTGAGAAT	:	181			
TrF3Ha40	:	-----CTACAGAGTTGTTG-ANCTTGTGAGNAT	:	28			
TrF3Ha41	:	-----	:	-			

FIGURE 75 (cont)

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      *           320           *           340           *           360
TrF3Ha1 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 353
TrF3Ha2 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 346
TrF3Ha3 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 359
TrF3Ha4 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 358
TrF3Ha5 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 318
TrF3Ha6 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 311
TrF3Ha7 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 313
TrF3Ha8 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 312
TrF3Ha9 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 309
TrF3Ha10 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 267
TrF3Ha11 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 306
TrF3Ha12 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 296
TrF3Ha13 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 304
TrF3Ha14 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 309
TrF3Ha15 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 311
TrF3Ha16 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 307
TrF3Ha17 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 309
TrF3Ha18 : ----- : -
TrF3Ha19 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 273
TrF3Ha20 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 306
TrF3Ha21 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 306
TrF3Ha22 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 302
TrF3Ha23 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 299
TrF3Ha24 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 297
TrF3Ha25 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 298
TrF3Ha26 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 299
TrF3Ha27 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 302
TrF3Ha28 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 302
TrF3Ha29 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 290
TrF3Ha30 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 295
TrF3Ha31 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 299
TrF3Ha32 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 296
TrF3Ha33 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 292
TrF3Ha34 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 289
TrF3Ha35 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 294
TrF3Ha36 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 293
TrF3Ha37 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 292
TrF3Ha38 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 292
TrF3Ha39 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 241
TrF3Ha40 : TGGGGTATTTTTCAGGTTGTTGATCATGGTGGTTGATACAAAACCTGTTTCTGAGATGACC : 88
TrF3Ha41 : ----- : -

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FIGURE 75 (cont)

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      *           380           *           400           *           420
TrF3Ha1 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :413
TrF3Ha2 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :406
TrF3Ha3 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :419
TrF3Ha4 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :418
TrF3Ha5 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :378
TrF3Ha6 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :371
TrF3Ha7 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :373
TrF3Ha8 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :372
TrF3Ha9 : CTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :369
TrF3Ha10 : ----- : -
TrF3Ha11 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :366
TrF3Ha12 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :356
TrF3Ha13 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :364
TrF3Ha14 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :369
TrF3Ha15 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :371
TrF3Ha16 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :367
TrF3Ha17 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :369
TrF3Ha18 : ----- : -
TrF3Ha19 : ----- : -
TrF3Ha20 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :366
TrF3Ha21 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :366
TrF3Ha22 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :362
TrF3Ha23 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :359
TrF3Ha24 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :357
TrF3Ha25 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :358
TrF3Ha26 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :359
TrF3Ha27 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :362
TrF3Ha28 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :362
TrF3Ha29 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :350
TrF3Ha30 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :355
TrF3Ha31 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :359
TrF3Ha32 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :356
TrF3Ha33 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :352
TrF3Ha34 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :349
TrF3Ha35 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :354
TrF3Ha36 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :353
TrF3Ha37 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :352
TrF3Ha38 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :352
TrF3Ha39 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :301
TrF3Ha40 : CGTTTTGCTAGAGAGTTTTTTTGCTTTGCCACCGGAAGAGAAGCTCCGGTTTGACATGTCC :148
TrF3Ha41 : ----- : -

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FIGURE 75 (cont)

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      *           440           *           460           *           480
TrF3Ha1 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :473
TrF3Ha2 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :466
TrF3Ha3 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :479
TrF3Ha4 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :478
TrF3Ha5 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :438
TrF3Ha6 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :431
TrF3Ha7 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :433
TrF3Ha8 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :432
TrF3Ha9 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :428
TrF3Ha10 : ----- : -
TrF3Ha11 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :426
TrF3Ha12 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :416
TrF3Ha13 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTNCAAGGAGAAAGCAGTGAAGGAT :424
TrF3Ha14 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :429
TrF3Ha15 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :431
TrF3Ha16 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :427
TrF3Ha17 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :429
TrF3Ha18 : ----- : -
TrF3Ha19 : ----- : -
TrF3Ha20 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :426
TrF3Ha21 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :426
TrF3Ha22 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :422
TrF3Ha23 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :419
TrF3Ha24 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :417
TrF3Ha25 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :418
TrF3Ha26 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :419
TrF3Ha27 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :422
TrF3Ha28 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAGCAGTGAAGGAT :422
TrF3Ha29 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :410
TrF3Ha30 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :415
TrF3Ha31 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :419
TrF3Ha32 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :416
TrF3Ha33 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTTCAAGGAGAAAGCAGTGAAGGAT :412
TrF3Ha34 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTNCAAGGAGAAAGCAGTGAAGGAT :409
TrF3Ha35 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :414
TrF3Ha36 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :413
TrF3Ha37 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :412
TrF3Ha38 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :412
TrF3Ha39 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :361
TrF3Ha40 : GGTGGTAAAAAGGGTGGTTTCATTGTCTCTAGTCATCTCCAAGGAGAAGCAGTGAAGGAT :208
TrF3Ha41 : ----- : -

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FIGURE 75 (cont)



	*	500	*	520	*	540
TrF3Ha1 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	533			
TrF3Ha2 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	526			
TrF3Ha3 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	539			
TrF3Ha4 :	TGCGAGAGAGCTNNNN-----	:	492			
TrF3Ha5 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	498			
TrF3Ha6 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	491			
TrF3Ha7 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	492			
TrF3Ha8 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	492			
TrF3Ha9 :	NNNNAAG-GNT-TTGGAANANNCNNNN-----	:	453			
TrF3Ha10 :	-----	:	-			
TrF3Ha11 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	486			
TrF3Ha12 :	TGGAGCGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	476			
TrF3Ha13 :	TGGAGAGAGCTNCNGCATATTTTTTN-----	:	450			
TrF3Ha14 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	489			
TrF3Ha15 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	491			
TrF3Ha16 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	487			
TrF3Ha17 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	489			
TrF3Ha18 :	-----	:	-			
TrF3Ha19 :	-----	:	-			
TrF3Ha20 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	486			
TrF3Ha21 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	486			
TrF3Ha22 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	482			
TrF3Ha23 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	479			
TrF3Ha24 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	477			
TrF3Ha25 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	478			
TrF3Ha26 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	479			
TrF3Ha27 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	482			
TrF3Ha28 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	482			
TrF3Ha29 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	470			
TrF3Ha30 :	TGN-----	:	418			
TrF3Ha31 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	479			
TrF3Ha32 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	476			
TrF3Ha33 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	472			
TrF3Ha34 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	469			
TrF3Ha35 :	TGGAGCGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	474			
TrF3Ha36 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	473			
TrF3Ha37 :	TGGAGCGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	472			
TrF3Ha38 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	472			
TrF3Ha39 :	TGGAGANAGCTAGTGCATATTTNTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	421			
TrF3Ha40 :	TGGAGAGAGCTAGTGCATATTTTTTCATACCCAATTAAACAAAGAGATTATTCAAGGTGG	:	268			
TrF3Ha41 :	-----	:	-			

**FIGURE 75 (cont)**

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	*	560	*	580	*	600	
TrF3Ha1	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCT	----				: 586
TrF3Ha2	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 586
TrF3Ha3	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 599
TrF3Ha4	:	-----					: -
TrF3Ha5	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 558
TrF3Ha6	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 551
TrF3Ha7	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 553
TrF3Ha8	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 552
TrF3Ha9	:	-----					: -
TrF3Ha10	:	-----					: -
TrF3Ha11	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 546
TrF3Ha12	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 536
TrF3Ha13	:	-----					: -
TrF3Ha14	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 549
TrF3Ha15	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 551
TrF3Ha16	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 547
TrF3Ha17	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 549
TrF3Ha18	:	-----					: -
TrF3Ha19	:	-----					: -
TrF3Ha20	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 546
TrF3Ha21	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 546
TrF3Ha22	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 542
TrF3Ha23	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 539
TrF3Ha24	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATG	---				: 535
TrF3Ha25	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 538
TrF3Ha26	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 539
TrF3Ha27	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 542
TrF3Ha28	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 542
TrF3Ha29	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 530
TrF3Ha30	:	-----					: -
TrF3Ha31	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 539
TrF3Ha32	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 536
TrF3Ha33	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 532
TrF3Ha34	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 529
TrF3Ha35	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 534
TrF3Ha36	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 533
TrF3Ha37	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 532
TrF3Ha38	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 532
TrF3Ha39	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 481
TrF3Ha40	:	CCAGACAAGCCAGAAGGATGGAAAGAGGTAACAGAAAAATACAGTGAAAACCTAATGAAT					: 328
TrF3Ha41	:	-----					: -

FIGURE 75 (cont)



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	*	620	*	640	*	660	
TrF3Ha1	:	-----		-----		-----	: -
TrF3Ha2	:	TTAGCTTGCAAG	-----				: 597
TrF3Ha3	:	TTAGCT	-----				: 605
TrF3Ha4	:	-----		-----		-----	: -
TrF3Ha5	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 618
TrF3Ha6	:	TTAGCTTGCAAG	CTATTGGAAG	-----			: 573
TrF3Ha7	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 613
TrF3Ha8	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 612
TrF3Ha9	:	-----		-----		-----	: -
TrF3Ha10	:	-----		-----		-----	: -
TrF3Ha11	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAG	-----		: 580
TrF3Ha12	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGA	TTAGAAAAN	-----	: 586
TrF3Ha13	:	-----		-----		-----	: -
TrF3Ha14	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 609
TrF3Ha15	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAG	-----		: 582
TrF3Ha16	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 607
TrF3Ha17	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	N	-----	: 593
TrF3Ha18	:	-----		-----		-----	: -
TrF3Ha19	:	-----		-----		-----	: -
TrF3Ha20	:	TTAGCTTGCAAG	CTATTGGAAGTTT	T	-----		: 572
TrF3Ha21	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATC	-----		: 575
TrF3Ha22	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAG	-----	: 596
TrF3Ha23	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 599
TrF3Ha24	:	-----		-----		-----	: -
TrF3Ha25	:	TTAGCTTGCAAAC	TATTGGAAN	-----			: 559
TrF3Ha26	:	TTAGCTTGCAAG	CTATTGGAAGTTT	T	-----		: 565
TrF3Ha27	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAN	-----	: 591
TrF3Ha28	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 602
TrF3Ha29	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 590
TrF3Ha30	:	-----		-----		-----	: -
TrF3Ha31	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAG	-----		: 570
TrF3Ha32	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 596
TrF3Ha33	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 592
TrF3Ha34	:	TTAGCT	-----				: 536
TrF3Ha35	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAAT	-----		: 572
TrF3Ha36	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGG	-----		: 573
TrF3Ha37	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGG	-----		: 573
TrF3Ha38	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGA	TTAGAAAAAG	-----	: 584
TrF3Ha39	:	TTAGCTTGCAAG	CTATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAG	-----	: 529
TrF3Ha40	:	TTAGCTTGCAAAC	TATTGGAAGTTT	TATCAGAAGCAATGGGTT	TAGAAAAAGAAGCTCTA		: 388
TrF3Ha41	:	-----		-----		-----	: -

FIGURE 75 (cont)

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		*	680	*	700	*	720	
TrF3Ha1	:	-----		-----		-----		:
TrF3Ha2	:	-----		-----		-----		:
TrF3Ha3	:	-----		-----		-----		:
TrF3Ha4	:	-----		-----		-----		:
TrF3Ha5	:	ACAAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT						: 678
TrF3Ha6	:	-----		-----		-----		:
TrF3Ha7	:	ACAAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT						: 673
TrF3Ha8	:	ACAAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT						: 672
TrF3Ha9	:	-----		-----		-----		:
TrF3Ha10	:	-----		-----		-----		:
TrF3Ha11	:	-----		-----		-----		:
TrF3Ha12	:	-----		-----		-----		:
TrF3Ha13	:	-----		-----		-----		:
TrF3Ha14	:	ACAAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT						: 669
TrF3Ha15	:	-----		-----		-----		:
TrF3Ha16	:	ACA-----		-----		-----		: 610
TrF3Ha17	:	-----		-----		-----		:
TrF3Ha18	:	-----		-----		-----		:
TrF3Ha19	:	-----		-----		-----		:
TrF3Ha20	:	-----		-----		-----		:
TrF3Ha21	:	-----		-----		-----		:
TrF3Ha22	:	-----		-----		-----		:
TrF3Ha23	:	ACAAAAGCATGTG-----		-----		-----		: 612
TrF3Ha24	:	-----		-----		-----		:
TrF3Ha25	:	-----		-----		-----		:
TrF3Ha26	:	-----		-----		-----		:
TrF3Ha27	:	-----		-----		-----		:
TrF3Ha28	:	ACAAAAG-----		-----		-----		: 609
TrF3Ha29	:	ACAAAAG-----		-----		-----		: 597
TrF3Ha30	:	-----		-----		-----		:
TrF3Ha31	:	-----		-----		-----		:
TrF3Ha32	:	ACAAAAGCATGT-----		-----		-----		: 608
TrF3Ha33	:	ACAAANNCNT-----		-----		-----		: 602
TrF3Ha34	:	-----		-----		-----		:
TrF3Ha35	:	-----		-----		-----		:
TrF3Ha36	:	-----		-----		-----		:
TrF3Ha37	:	-----		-----		-----		:
TrF3Ha38	:	-----		-----		-----		:
TrF3Ha39	:	-----		-----		-----		:
TrF3Ha40	:	ACAAAAGCATGTGTTGATATGGATCAAAAAGTTGTTATAAATTATTACCCAAAATGCCCT						: 448
TrF3Ha41	:	-----ATCMAAAAAGTTGTCATAAANTATTACCCMAAAATGCCCT						: 38

## FIGURE 75 (cont)

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	*	740	*	760	*	780		
TrF3Ha1	:	-----		-----		-----	:	-
TrF3Ha2	:	-----		-----		-----	:	-
TrF3Ha3	:	-----		-----		-----	:	-
TrF3Ha4	:	-----		-----		-----	:	-
TrF3Ha5	:	GAACCTGACCTT		-----		-----	:	690
TrF3Ha6	:	-----		-----		-----	:	-
TrF3Ha7	:	GAACCTGACCTCGCACTT		GCCGTAAACGNCACACTGACCCNGA		-----	:	716
TrF3Ha8	:	GAACCTGACCTCACACTTGGCCTTAAACGTCACACTGACCCTGGCACAN		-----		-----	:	721
TrF3Ha9	:	-----		-----		-----	:	-
TrF3Ha10	:	-----		-----		-----	:	-
TrF3Ha11	:	-----		-----		-----	:	-
TrF3Ha12	:	-----		-----		-----	:	-
TrF3Ha13	:	-----		-----		-----	:	-
TrF3Ha14	:	GAACCTGACCTC		-----		-----	:	681
TrF3Ha15	:	-----		-----		-----	:	-
TrF3Ha16	:	-----		-----		-----	:	-
TrF3Ha17	:	-----		-----		-----	:	-
TrF3Ha18	:	-----		-----		-----	:	-
TrF3Ha19	:	-----		-----		-----	:	-
TrF3Ha20	:	-----		-----		-----	:	-
TrF3Ha21	:	-----		-----		-----	:	-
TrF3Ha22	:	-----		-----		-----	:	-
TrF3Ha23	:	-----		-----		-----	:	-
TrF3Ha24	:	-----		-----		-----	:	-
TrF3Ha25	:	-----		-----		-----	:	-
TrF3Ha26	:	-----		-----		-----	:	-
TrF3Ha27	:	-----		-----		-----	:	-
TrF3Ha28	:	-----		-----		-----	:	-
TrF3Ha29	:	-----		-----		-----	:	-
TrF3Ha30	:	-----		-----		-----	:	-
TrF3Ha31	:	-----		-----		-----	:	-
TrF3Ha32	:	-----		-----		-----	:	-
TrF3Ha33	:	-----		-----		-----	:	-
TrF3Ha34	:	-----		-----		-----	:	-
TrF3Ha35	:	-----		-----		-----	:	-
TrF3Ha36	:	-----		-----		-----	:	-
TrF3Ha37	:	-----		-----		-----	:	-
TrF3Ha38	:	-----		-----		-----	:	-
TrF3Ha39	:	-----		-----		-----	:	-
TrF3Ha40	:	GAACCTGACCTCACACTTGGCCTTAAACGTCACACTGACCCTGGCACAATTACTCTTTTG		-----		-----	:	508
TrF3Ha41	:	GAACCTGACCTCACACTTGGCCTTAAACGTCACACTGACCCTGGNACAATTACTCTTTTG		-----		-----	:	98

FIGURE 75 (cont)

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		*	800	*	820	*	840	
TrF3Ha1	:	-----		-----		-----		:
TrF3Ha2	:	-----		-----		-----		:
TrF3Ha3	:	-----		-----		-----		:
TrF3Ha4	:	-----		-----		-----		:
TrF3Ha5	:	-----		-----		-----		:
TrF3Ha6	:	-----		-----		-----		:
TrF3Ha7	:	-----		-----		-----		:
TrF3Ha8	:	-----		-----		-----		:
TrF3Ha9	:	-----		-----		-----		:
TrF3Ha10	:	-----		-----		-----		:
TrF3Ha11	:	-----		-----		-----		:
TrF3Ha12	:	-----		-----		-----		:
TrF3Ha13	:	-----		-----		-----		:
TrF3Ha14	:	-----		-----		-----		:
TrF3Ha15	:	-----		-----		-----		:
TrF3Ha16	:	-----		-----		-----		:
TrF3Ha17	:	-----		-----		-----		:
TrF3Ha18	:	-----		-----		-----		:
TrF3Ha19	:	-----		-----		-----		:
TrF3Ha20	:	-----		-----		-----		:
TrF3Ha21	:	-----		-----		-----		:
TrF3Ha22	:	-----		-----		-----		:
TrF3Ha23	:	-----		-----		-----		:
TrF3Ha24	:	-----		-----		-----		:
TrF3Ha25	:	-----		-----		-----		:
TrF3Ha26	:	-----		-----		-----		:
TrF3Ha27	:	-----		-----		-----		:
TrF3Ha28	:	-----		-----		-----		:
TrF3Ha29	:	-----		-----		-----		:
TrF3Ha30	:	-----		-----		-----		:
TrF3Ha31	:	-----		-----		-----		:
TrF3Ha32	:	-----		-----		-----		:
TrF3Ha33	:	-----		-----		-----		:
TrF3Ha34	:	-----		-----		-----		:
TrF3Ha35	:	-----		-----		-----		:
TrF3Ha36	:	-----		-----		-----		:
TrF3Ha37	:	-----		-----		-----		:
TrF3Ha38	:	-----		-----		-----		:
TrF3Ha39	:	-----		-----		-----		:
TrF3Ha40	:	CTTCAAGATCAAGTTGGTGGGCTTCAAGCTACCAAAGATAATGGTAAGACGTGGATTACA						: 568
TrF3Ha41	:	CTTCAAGATCAAGTTGGTGGCCTTCAAGCTACCAAAGATAATCGTAAGACGTGGATTACA						: 158

## FIGURE 75 (cont)

# 135/271

		*	860	*	880	*	900	
TrF3Ha1	:	-----		-----		-----		:
TrF3Ha2	:	-----		-----		-----		:
TrF3Ha3	:	-----		-----		-----		:
TrF3Ha4	:	-----		-----		-----		:
TrF3Ha5	:	-----		-----		-----		:
TrF3Ha6	:	-----		-----		-----		:
TrF3Ha7	:	-----		-----		-----		:
TrF3Ha8	:	-----		-----		-----		:
TrF3Ha9	:	-----		-----		-----		:
TrF3Ha10	:	-----		-----		-----		:
TrF3Ha11	:	-----		-----		-----		:
TrF3Ha12	:	-----		-----		-----		:
TrF3Ha13	:	-----		-----		-----		:
TrF3Ha14	:	-----		-----		-----		:
TrF3Ha15	:	-----		-----		-----		:
TrF3Ha16	:	-----		-----		-----		:
TrF3Ha17	:	-----		-----		-----		:
TrF3Ha18	:	-----		-----		-----		:
TrF3Ha19	:	-----		-----		-----		:
TrF3Ha20	:	-----		-----		-----		:
TrF3Ha21	:	-----		-----		-----		:
TrF3Ha22	:	-----		-----		-----		:
TrF3Ha23	:	-----		-----		-----		:
TrF3Ha24	:	-----		-----		-----		:
TrF3Ha25	:	-----		-----		-----		:
TrF3Ha26	:	-----		-----		-----		:
TrF3Ha27	:	-----		-----		-----		:
TrF3Ha28	:	-----		-----		-----		:
TrF3Ha29	:	-----		-----		-----		:
TrF3Ha30	:	-----		-----		-----		:
TrF3Ha31	:	-----		-----		-----		:
TrF3Ha32	:	-----		-----		-----		:
TrF3Ha33	:	-----		-----		-----		:
TrF3Ha34	:	-----		-----		-----		:
TrF3Ha35	:	-----		-----		-----		:
TrF3Ha36	:	-----		-----		-----		:
TrF3Ha37	:	-----		-----		-----		:
TrF3Ha38	:	-----		-----		-----		:
TrF3Ha39	:	-----		-----		-----		:
TrF3Ha40	:	GTTCAACCAGTTGAAGGTGCTTTTGTGTTAATCTTGGAGACCATGGTCACTATCTAAGT						: 628
TrF3Ha41	:	GTTCAACCAGTTGAAGGTGCTTTTGTGTTAATCTTGGAGACCATGGTCACTATCTAAGT						: 218

## FIGURE 75 (cont)

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	*	920	*	940	*	960	
TrF3Ha1	:	-----	:	-----	:	-----	:
TrF3Ha2	:	-----	:	-----	:	-----	:
TrF3Ha3	:	-----	:	-----	:	-----	:
TrF3Ha4	:	-----	:	-----	:	-----	:
TrF3Ha5	:	-----	:	-----	:	-----	:
TrF3Ha6	:	-----	:	-----	:	-----	:
TrF3Ha7	:	-----	:	-----	:	-----	:
TrF3Ha8	:	-----	:	-----	:	-----	:
TrF3Ha9	:	-----	:	-----	:	-----	:
TrF3Ha10	:	-----	:	-----	:	-----	:
TrF3Ha11	:	-----	:	-----	:	-----	:
TrF3Ha12	:	-----	:	-----	:	-----	:
TrF3Ha13	:	-----	:	-----	:	-----	:
TrF3Ha14	:	-----	:	-----	:	-----	:
TrF3Ha15	:	-----	:	-----	:	-----	:
TrF3Ha16	:	-----	:	-----	:	-----	:
TrF3Ha17	:	-----	:	-----	:	-----	:
TrF3Ha18	:	-----	:	-----	:	-----	:
TrF3Ha19	:	-----	:	-----	:	-----	:
TrF3Ha20	:	-----	:	-----	:	-----	:
TrF3Ha21	:	-----	:	-----	:	-----	:
TrF3Ha22	:	-----	:	-----	:	-----	:
TrF3Ha23	:	-----	:	-----	:	-----	:
TrF3Ha24	:	-----	:	-----	:	-----	:
TrF3Ha25	:	-----	:	-----	:	-----	:
TrF3Ha26	:	-----	:	-----	:	-----	:
TrF3Ha27	:	-----	:	-----	:	-----	:
TrF3Ha28	:	-----	:	-----	:	-----	:
TrF3Ha29	:	-----	:	-----	:	-----	:
TrF3Ha30	:	-----	:	-----	:	-----	:
TrF3Ha31	:	-----	:	-----	:	-----	:
TrF3Ha32	:	-----	:	-----	:	-----	:
TrF3Ha33	:	-----	:	-----	:	-----	:
TrF3Ha34	:	-----	:	-----	:	-----	:
TrF3Ha35	:	-----	:	-----	:	-----	:
TrF3Ha36	:	-----	:	-----	:	-----	:
TrF3Ha37	:	-----	:	-----	:	-----	:
TrF3Ha38	:	-----	:	-----	:	-----	:
TrF3Ha39	:	-----	:	-----	:	-----	:
TrF3Ha40	:	AATGGACGGTTCAAAAATGCTGACCAATCAAGCAGTGGTGAATTTCGAACTACAGCCGNTTA					: 688
TrF3Ha41	:	AATGGACGGTTCAAAAATGCTGACCACCAAGCAGTGGTGAATTTCGAACTACAGCCGTTTA					: 278

## FIGURE 75 (cont)

# 137/271

	*	980	*	1000	*	1020		
TrF3Ha1	:	-----		-----		-----	:	-
TrF3Ha2	:	-----		-----		-----	:	-
TrF3Ha3	:	-----		-----		-----	:	-
TrF3Ha4	:	-----		-----		-----	:	-
TrF3Ha5	:	-----		-----		-----	:	-
TrF3Ha6	:	-----		-----		-----	:	-
TrF3Ha7	:	-----		-----		-----	:	-
TrF3Ha8	:	-----		-----		-----	:	-
TrF3Ha9	:	-----		-----		-----	:	-
TrF3Ha10	:	-----		-----		-----	:	-
TrF3Ha11	:	-----		-----		-----	:	-
TrF3Ha12	:	-----		-----		-----	:	-
TrF3Ha13	:	-----		-----		-----	:	-
TrF3Ha14	:	-----		-----		-----	:	-
TrF3Ha15	:	-----		-----		-----	:	-
TrF3Ha16	:	-----		-----		-----	:	-
TrF3Ha17	:	-----		-----		-----	:	-
TrF3Ha18	:	-----		-----		-----	:	-
TrF3Ha19	:	-----		-----		-----	:	-
TrF3Ha20	:	-----		-----		-----	:	-
TrF3Ha21	:	-----		-----		-----	:	-
TrF3Ha22	:	-----		-----		-----	:	-
TrF3Ha23	:	-----		-----		-----	:	-
TrF3Ha24	:	-----		-----		-----	:	-
TrF3Ha25	:	-----		-----		-----	:	-
TrF3Ha26	:	-----		-----		-----	:	-
TrF3Ha27	:	-----		-----		-----	:	-
TrF3Ha28	:	-----		-----		-----	:	-
TrF3Ha29	:	-----		-----		-----	:	-
TrF3Ha30	:	-----		-----		-----	:	-
TrF3Ha31	:	-----		-----		-----	:	-
TrF3Ha32	:	-----		-----		-----	:	-
TrF3Ha33	:	-----		-----		-----	:	-
TrF3Ha34	:	-----		-----		-----	:	-
TrF3Ha35	:	-----		-----		-----	:	-
TrF3Ha36	:	-----		-----		-----	:	-
TrF3Ha37	:	-----		-----		-----	:	-
TrF3Ha38	:	-----		-----		-----	:	-
TrF3Ha39	:	-----		-----		-----	:	-
TrF3Ha40	:	TCAATAGCAA		-----		-----	:	698
TrF3Ha41	:	TCAATAGCAACATTTCAAAATCCAGCTCCCGATGCAACTGTATACCCTTTGAAGATTAGA		-----		-----	:	338

## FIGURE 75 (cont)

# 138/271

	*	1040	*	1060	*	1080	
TrF3Ha1	:	-----		-----		-----	:
TrF3Ha2	:	-----		-----		-----	:
TrF3Ha3	:	-----		-----		-----	:
TrF3Ha4	:	-----		-----		-----	:
TrF3Ha5	:	-----		-----		-----	:
TrF3Ha6	:	-----		-----		-----	:
TrF3Ha7	:	-----		-----		-----	:
TrF3Ha8	:	-----		-----		-----	:
TrF3Ha9	:	-----		-----		-----	:
TrF3Ha10	:	-----		-----		-----	:
TrF3Ha11	:	-----		-----		-----	:
TrF3Ha12	:	-----		-----		-----	:
TrF3Ha13	:	-----		-----		-----	:
TrF3Ha14	:	-----		-----		-----	:
TrF3Ha15	:	-----		-----		-----	:
TrF3Ha16	:	-----		-----		-----	:
TrF3Ha17	:	-----		-----		-----	:
TrF3Ha18	:	-----		-----		-----	:
TrF3Ha19	:	-----		-----		-----	:
TrF3Ha20	:	-----		-----		-----	:
TrF3Ha21	:	-----		-----		-----	:
TrF3Ha22	:	-----		-----		-----	:
TrF3Ha23	:	-----		-----		-----	:
TrF3Ha24	:	-----		-----		-----	:
TrF3Ha25	:	-----		-----		-----	:
TrF3Ha26	:	-----		-----		-----	:
TrF3Ha27	:	-----		-----		-----	:
TrF3Ha28	:	-----		-----		-----	:
TrF3Ha29	:	-----		-----		-----	:
TrF3Ha30	:	-----		-----		-----	:
TrF3Ha31	:	-----		-----		-----	:
TrF3Ha32	:	-----		-----		-----	:
TrF3Ha33	:	-----		-----		-----	:
TrF3Ha34	:	-----		-----		-----	:
TrF3Ha35	:	-----		-----		-----	:
TrF3Ha36	:	-----		-----		-----	:
TrF3Ha37	:	-----		-----		-----	:
TrF3Ha38	:	-----		-----		-----	:
TrF3Ha39	:	-----		-----		-----	:
TrF3Ha40	:	-----		-----		-----	:
TrF3Ha41	:	GAGGGTGAAAAATCTGTGTTGGAAGAACCAATCACTTTTGCTGAAATGTATAGAAGGAAG					:398

## FIGURE 75 (cont)



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		*	1100	*	1120	*	1140	
TrF3Ha1	:	-----		-----		-----		:
TrF3Ha2	:	-----		-----		-----		:
TrF3Ha3	:	-----		-----		-----		:
TrF3Ha4	:	-----		-----		-----		:
TrF3Ha5	:	-----		-----		-----		:
TrF3Ha6	:	-----		-----		-----		:
TrF3Ha7	:	-----		-----		-----		:
TrF3Ha8	:	-----		-----		-----		:
TrF3Ha9	:	-----		-----		-----		:
TrF3Ha10	:	-----		-----		-----		:
TrF3Ha11	:	-----		-----		-----		:
TrF3Ha12	:	-----		-----		-----		:
TrF3Ha13	:	-----		-----		-----		:
TrF3Ha14	:	-----		-----		-----		:
TrF3Ha15	:	-----		-----		-----		:
TrF3Ha16	:	-----		-----		-----		:
TrF3Ha17	:	-----		-----		-----		:
TrF3Ha18	:	-----		-----		-----		:
TrF3Ha19	:	-----		-----		-----		:
TrF3Ha20	:	-----		-----		-----		:
TrF3Ha21	:	-----		-----		-----		:
TrF3Ha22	:	-----		-----		-----		:
TrF3Ha23	:	-----		-----		-----		:
TrF3Ha24	:	-----		-----		-----		:
TrF3Ha25	:	-----		-----		-----		:
TrF3Ha26	:	-----		-----		-----		:
TrF3Ha27	:	-----		-----		-----		:
TrF3Ha28	:	-----		-----		-----		:
TrF3Ha29	:	-----		-----		-----		:
TrF3Ha30	:	-----		-----		-----		:
TrF3Ha31	:	-----		-----		-----		:
TrF3Ha32	:	-----		-----		-----		:
TrF3Ha33	:	-----		-----		-----		:
TrF3Ha34	:	-----		-----		-----		:
TrF3Ha35	:	-----		-----		-----		:
TrF3Ha36	:	-----		-----		-----		:
TrF3Ha37	:	-----		-----		-----		:
TrF3Ha38	:	-----		-----		-----		:
TrF3Ha39	:	-----		-----		-----		:
TrF3Ha40	:	-----		-----		-----		:
TrF3Ha41	:	ATGACCAAAGACCTTGAAATTGCTAGGATGAAGAAGTTGGCTAAGGAACAACAACCTTAGG						: 458

FIGURE 75 (cont)

# 140/271

	*	1160	*	1180	*	1200	
TrF3Ha1	:	-----	-----	-----	-----	-----	:
TrF3Ha2	:	-----	-----	-----	-----	-----	:
TrF3Ha3	:	-----	-----	-----	-----	-----	:
TrF3Ha4	:	-----	-----	-----	-----	-----	:
TrF3Ha5	:	-----	-----	-----	-----	-----	:
TrF3Ha6	:	-----	-----	-----	-----	-----	:
TrF3Ha7	:	-----	-----	-----	-----	-----	:
TrF3Ha8	:	-----	-----	-----	-----	-----	:
TrF3Ha9	:	-----	-----	-----	-----	-----	:
TrF3Ha10	:	-----	-----	-----	-----	-----	:
TrF3Ha11	:	-----	-----	-----	-----	-----	:
TrF3Ha12	:	-----	-----	-----	-----	-----	:
TrF3Ha13	:	-----	-----	-----	-----	-----	:
TrF3Ha14	:	-----	-----	-----	-----	-----	:
TrF3Ha15	:	-----	-----	-----	-----	-----	:
TrF3Ha16	:	-----	-----	-----	-----	-----	:
TrF3Ha17	:	-----	-----	-----	-----	-----	:
TrF3Ha18	:	-----	-----	-----	-----	-----	:
TrF3Ha19	:	-----	-----	-----	-----	-----	:
TrF3Ha20	:	-----	-----	-----	-----	-----	:
TrF3Ha21	:	-----	-----	-----	-----	-----	:
TrF3Ha22	:	-----	-----	-----	-----	-----	:
TrF3Ha23	:	-----	-----	-----	-----	-----	:
TrF3Ha24	:	-----	-----	-----	-----	-----	:
TrF3Ha25	:	-----	-----	-----	-----	-----	:
TrF3Ha26	:	-----	-----	-----	-----	-----	:
TrF3Ha27	:	-----	-----	-----	-----	-----	:
TrF3Ha28	:	-----	-----	-----	-----	-----	:
TrF3Ha29	:	-----	-----	-----	-----	-----	:
TrF3Ha30	:	-----	-----	-----	-----	-----	:
TrF3Ha31	:	-----	-----	-----	-----	-----	:
TrF3Ha32	:	-----	-----	-----	-----	-----	:
TrF3Ha33	:	-----	-----	-----	-----	-----	:
TrF3Ha34	:	-----	-----	-----	-----	-----	:
TrF3Ha35	:	-----	-----	-----	-----	-----	:
TrF3Ha36	:	-----	-----	-----	-----	-----	:
TrF3Ha37	:	-----	-----	-----	-----	-----	:
TrF3Ha38	:	-----	-----	-----	-----	-----	:
TrF3Ha39	:	-----	-----	-----	-----	-----	:
TrF3Ha40	:	-----	-----	-----	-----	-----	:
TrF3Ha41	:	GACTTGGAGGAGAACAAGACTAAATATGAGGCCAAACCTTTGAATGAGATCTTTGCTTAA					:

: 518

## FIGURE 75 (cont)

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	*	1220	*	1240	*	1260	
TrF3Ha1	:	-----		-----		-----	:
TrF3Ha2	:	-----		-----		-----	:
TrF3Ha3	:	-----		-----		-----	:
TrF3Ha4	:	-----		-----		-----	:
TrF3Ha5	:	-----		-----		-----	:
TrF3Ha6	:	-----		-----		-----	:
TrF3Ha7	:	-----		-----		-----	:
TrF3Ha8	:	-----		-----		-----	:
TrF3Ha9	:	-----		-----		-----	:
TrF3Ha10	:	-----		-----		-----	:
TrF3Ha11	:	-----		-----		-----	:
TrF3Ha12	:	-----		-----		-----	:
TrF3Ha13	:	-----		-----		-----	:
TrF3Ha14	:	-----		-----		-----	:
TrF3Ha15	:	-----		-----		-----	:
TrF3Ha16	:	-----		-----		-----	:
TrF3Ha17	:	-----		-----		-----	:
TrF3Ha18	:	-----		-----		-----	:
TrF3Ha19	:	-----		-----		-----	:
TrF3Ha20	:	-----		-----		-----	:
TrF3Ha21	:	-----		-----		-----	:
TrF3Ha22	:	-----		-----		-----	:
TrF3Ha23	:	-----		-----		-----	:
TrF3Ha24	:	-----		-----		-----	:
TrF3Ha25	:	-----		-----		-----	:
TrF3Ha26	:	-----		-----		-----	:
TrF3Ha27	:	-----		-----		-----	:
TrF3Ha28	:	-----		-----		-----	:
TrF3Ha29	:	-----		-----		-----	:
TrF3Ha30	:	-----		-----		-----	:
TrF3Ha31	:	-----		-----		-----	:
TrF3Ha32	:	-----		-----		-----	:
TrF3Ha33	:	-----		-----		-----	:
TrF3Ha34	:	-----		-----		-----	:
TrF3Ha35	:	-----		-----		-----	:
TrF3Ha36	:	-----		-----		-----	:
TrF3Ha37	:	-----		-----		-----	:
TrF3Ha38	:	-----		-----		-----	:
TrF3Ha39	:	-----		-----		-----	:
TrF3Ha40	:	-----		-----		-----	:
TrF3Ha41	:	TTAATTAGTCTTAATT	TAAATAATTAATAAATTT	TAGACTTAATTTACATATAATAATTT			: 578

## FIGURE 75 (cont)

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TrF3Ha1	:	-	:	-
TrF3Ha2	:	-	:	-
TrF3Ha3	:	-	:	-
TrF3Ha4	:	-	:	-
TrF3Ha5	:	-	:	-
TrF3Ha6	:	-	:	-
TrF3Ha7	:	-	:	-
TrF3Ha8	:	-	:	-
TrF3Ha9	:	-	:	-
TrF3Ha10	:	-	:	-
TrF3Ha11	:	-	:	-
TrF3Ha12	:	-	:	-
TrF3Ha13	:	-	:	-
TrF3Ha14	:	-	:	-
TrF3Ha15	:	-	:	-
TrF3Ha16	:	-	:	-
TrF3Ha17	:	-	:	-
TrF3Ha18	:	-	:	-
TrF3Ha19	:	-	:	-
TrF3Ha20	:	-	:	-
TrF3Ha21	:	-	:	-
TrF3Ha22	:	-	:	-
TrF3Ha23	:	-	:	-
TrF3Ha24	:	-	:	-
TrF3Ha25	:	-	:	-
TrF3Ha26	:	-	:	-
TrF3Ha27	:	-	:	-
TrF3Ha28	:	-	:	-
TrF3Ha29	:	-	:	-
TrF3Ha30	:	-	:	-
TrF3Ha31	:	-	:	-
TrF3Ha32	:	-	:	-
TrF3Ha33	:	-	:	-
TrF3Ha34	:	-	:	-
TrF3Ha35	:	-	:	-
TrF3Ha36	:	-	:	-
TrF3Ha37	:	-	:	-
TrF3Ha38	:	-	:	-
TrF3Ha39	:	-	:	-
TrF3Ha40	:	-	:	-
TrF3Ha41	:	T	:	579

## FIGURE 75 (cont)

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```

      *           20           *           40           *           60
TrF3Hb : GNAGCATAACATAAACCCCTGTNCCCGATTNATGTAACACAATCTCCCCTTTTCTTATTAC : 60

      *           80           *           100          *           120
TrF3Hb : AAGTAAAATACCATAACACAATAATATGAATACCATAATCTTGAATCATACAAACAACCT : 120

      *           140          *           160          *           180
TrF3Hb : TGGATCAAACAAAACAACAACCATGGTTGATCTAGAAACAGAACCAAGTTCACCATTTAT : 180

      *           200          *           220          *           240
TrF3Hb : TCAATCCCCAGAACACAGACCAAAATCCTCAATAATCATTGCTGAAGGTATCCCTCTAAT : 240

      *           260          *           280          *           300
TrF3Hb : TGATCTCACTCCTATAAACTACAAAGATGAAATCATCACCAACCCACTTTCCATTGAAGA : 300

      *           320          *           340          *           360
TrF3Hb : CTTAGTCAAAGAAATAGGCAAAGCATGTAAAGAATGGGGTTTCTTTCAAGTGATTAATCA : 360

      *           380          *           400          *           420
TrF3Hb : CAAAGTTCCTTTGGATAAACGTGAAAGGATTGAAGAATCTTCAAAGAAGTTTTTTGAACT : 420

      *           440          *           460          *           480
TrF3Hb : TAGTTTGGAGGAAAAACTTAAGGTGAGAAGAGATGAAGTTAATTTGCTTGGTTATTTTGA : 480

      *           500          *           520          *           540
TrF3Hb : AGCTGAGCATACAAAAAATGTTAGGGACTGGAAGGAAATTTATGATTTTAATGTGCAACA : 540

      *           560          *           580          *           600
TrF3Hb : ACCAACTTTTATACCACCTTCGGATGACCAAAGTTTTTCAGTTTCAATGGGAAAATCGATG : 600

TrF3Hb : G : 601

```

## FIGURE 76

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TrF3Hb : MNTIILNHTNNLGSNKTTTMVDLETEPSSPFIQSPEHRPKSSIIIAEGIPLIDLTPINYK : 60

TrF3Hb : DEIITNPLSIEDLVKEIGKACKEWGFFQVINHKVPLDKRERIEESSKKFFELSLEEKLV : 120

TrF3Hb : RRDEVNLLGYFEAEHTKNVRDWKEIYDFNVQQPTFIPPSDDQSFQFQWENRW : 172

## FIGURE 77

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```

      *           20           *           40           *           60
TrF3Hc : TTACCCAACAATNATGTGTGACTGATGTTAGTGTACCAGGAAAGATGGGAGAGGTGGATC : 60

      *           80           *           100          *           120
TrF3Hc : CAGCTTCTCTCAAATCCAGAAAATAGGCCAAAAC TTCCATAATCCAAGCTGAAGGAATT : 120

      *           140          *           160          *           180
TrF3Hc : CCTGTAATCAATCTCTCCCCATTAATTCACCACACAGTTCAAGACTCCTCTGCCATTGAA : 180

      *           200          *           220          *           240
TrF3Hc : AGCTTAGTCAAAGAAATAGGAAATGCTTGCAAGGAATGGGGTTTCTTCCAAGTAACAAAC : 240

      *           260          *           280          *           300
TrF3Hc : CATGGTGTCCCTCTAAATCTAAGGCTCAGACTCGAGGAAGCTACCAAAGTTTCTTTTGCA : 300

      *           320          *           340          *           360
TrF3Hc : CAGAGTTTGGAGGAGAAGAGGAAGCTTACCGTAGATGATAACAGTTTGCCTGGTTATCAT : 360

      *           380          *           400          *           420
TrF3Hc : GATACAGAGCACACCAAGAATGTCAGAGACTGGAAAGAAGTGTGATTTTTTATCCAAA : 420

      *           440          *           460          *           480
TrF3Hc : GACCCCACTTTGATTCTCTGAATTCTGATGAACATGATGATCGAGTCACTCAATGGACT : 480

      *           500          *           520          *           540
TrF3Hc : AATCCATCCCCTCAATATCCTCCAAACTTCAAAGTTATTTTGGAAGAGTATATTAAAGAG : 540

      *           560          *           580
TrF3Hc : ATGGAAGCTAGGCTTTAAGTTGCTAGAGCTTATAGCTTTGAGC : 585

```

## FIGURE 78

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TrF3Hc :                   \*                  20                  \*                  40                  \*                  60  
          : MLVYQERWERWIQLSSNPENRPKLSIIQAEGIPVINLSPLIHHTVQDSSAIESLVKEIGN : 60

TrF3Hc :                   \*                  80                  \*                  100                  \*                  120  
          : ACKEWGFFQVTNHGVPLNLRRLRLLEEATKVFFAQSLLEKRKLTVDDNSLPGYHDEHTKNV : 120

TrF3Hc :                   \*                  140                  \*                  160                  \*                  180  
          : RDWKEVFDLFLSKDPTLIPLNSDEHDDRVTQWTNPSPQYPPNFKVILEEYIKEMEKLGFKL : 180

TrF3Hc : LELIALS : 187

## FIGURE 79



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TrF3'Ha : GGAATGGTGGAGGCGAATGTGACCCTAGGGCTGATGAATTAGTAATGGTAGTTGAGCTT : 60

TrF3'Ha : ATGGCGTTAGCTGGAGTTTCAATATTGGTGATTTTGTTCCTGCTTTGGAATGGTTAGAT :120

TrF3'Ha : ATTCAAGGTGTACAAGGAAAAATGAAGAAATTACATAAAAGATTGATGCATTTTAACT :180

TrF3'Ha : AGCATTATTGAAGATCACATGATTTCCAAGAGTGAGAAGCATAATGACTTATTGAGTACG :240

TrF3'Ha : TTGTTATCACTAAAAGAAAAAGTTGATGAGGATGGTGACAACTTAATGATACTGAGATC :300

TrF3'Ha : AAAGCATTACTCTTGAACATGTTTCACAGCTGGAACAGACACATCATCAAGCACACAGAG :360

TrF3'Ha : TGGGCTATTGCTGAACTAATAAAAAATCCAAACTAATGATTCGTGTTCAAATGAGTTG :420

TrF3'Ha : GACACTGTTGTGGGCCGAGACAAGCTTGTAAGTGAACAAGACTTGGCCCATCTTCCTTAC :480

TrF3'Ha : TTAGAGGCTGTAATAAAGGAGACATTTTCGTCTCCATCCATCAACCCCTCTTCTCTCCCA :540

TrF3'Ha : CGTGTGCAACAAATAGTTGTGAAATCCTCGACTATCACATTCCCAAAGGTGCAACTCTC :600

TrF3'Ha : TTGG : 604

## FIGURE 80

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```

      *           20           *           40           *           60
TrF3'Ha : GNGGGECDPRADELVMVVELMALAGVFNIGDFVPALWLDIQGVQGMKKLHKRFDAFLT : 60

      *           80           *           100          *           120
TrF3'Ha : SIIEDHMISKSEKHNDLLSTLLSLKEKVDEDGDKLNDTEIKALLNMFTAGTDTSSSTTE :120

      *           140          *           160          *           180
TrF3'Ha : WAIAELIKNPCLMIRVQNELDTVVGRDKLVTEQDLAHLPLYEAVIKETFRLHPSTPLSLP :180

      *           200
TrF3'Ha : RVATNSCEILDYHIPKGATLL :201
```

## FIGURE 81

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```

      *           20           *           40           *           60
TrF3'Ha1 : GGG AATGGTGGAGGCGAATGTGACCCTAGGGCTGATGAATTTAAGTAATGGTAGTTGAGC : 60
TrF3'Ha2 : ---AATGGTGGAGGCGAATGTGACCCTAGGGCTGATGAATTTAAGTAATGGTAGTTGAGC : 57

      *           80           *           100          *           120
TrF3'Ha1 : TTATGGCGTTAGCTGGAGTTTTCAATATTGGTGATTTTGTTCCTGCTTTGGAATGGTTAG : 120
TrF3'Ha2 : TTATGGCGTTAGCTGGAGTTTTCAATATTGGTGATTTTGTTCCTGCTTTGGAATGGTTAG : 117

      *           140          *           160           *           180
TrF3'Ha1 : ATATTCAAGGTGTACAAGGAAAAATGAAGAAATTACATAAAAGATTTGATGCATTTTAA : 180
TrF3'Ha2 : ATATTCAAGGTGTACAAGGAAAAATGAAGAAATTACATAAAAGATTTGATGCATTTTAA : 177

      *           200          *           220           *           240
TrF3'Ha1 : CTAGCATTATTGAAGATCACATGATTTCCAAGAGTGAGAAGCATAATGACTTATTGAGTA : 240
TrF3'Ha2 : CTAGCATTATTGAAGATCACATGATTTCCAAGAGTGAGAAGCATAATGACTTATTGAGTA : 237

      *           260          *           280           *           300
TrF3'Ha1 : CGTTGTTATCACTAAAAGAAAAAGTTGATGAGGATGGTGACAACTTAATGATACTGAGA : 300
TrF3'Ha2 : CGTTGTTATCACTAAAAGAAAAAGTTGATGAGGATGGTGACAACTTAATGATACTGAGA : 297

      *           320          *           340           *           360
TrF3'Ha1 : TCAAAGCATTACTCTTGAACATGTTTCACAGCTGGAACAGACACATCATCAAGCACAACAG : 360
TrF3'Ha2 : TCAAAGCATTACTCTTGAACATGTTTCACAGCTGGAACAGACACATCATCAAGCACAACAG : 357

      *           380          *           400           *           420
TrF3'Ha1 : AGTGGGCTATTGCTGAACTAATAAAAAATCCAAAACATAATGATTCGTGTTCAAAATGAGT : 420
TrF3'Ha2 : AGTGGGCTATTGCTGAACTAATAAAAAATCCAAAACATAATGATTCGTGTTCAAAATGAGT : 417

      *           440          *           460           *           480
TrF3'Ha1 : TGGACACTGTTGTGGGCCGAGACAAGCTTGTAACCTGAACAAGACTTGGCCCATCTTCCTT : 480
TrF3'Ha2 : TGGACACTGTTGTGGGCCGAGACAAGCTTGTAACCTGAACAAGACTTGGCCCATCTTCCTT : 477

      *           500          *           520           *           540
TrF3'Ha1 : ACTTAGAGGCTGTAATAAAGGAGACATTTTCGTCTCCATCCATCAACCCCTCTTTCTCTCC : 540
TrF3'Ha2 : ACTTAGAGGCTGTAATAAAGGAGACATTTTCGTCTCCATCCATCAACCCCTCTTTCTCTCC : 537

      *           560          *           580           *           600
TrF3'Ha1 : CACGTGTTGCAACAAATAGTTGTGAAATCCTCGACTATCAC----- : 581
TrF3'Ha2 : CACGTGTTGCAACAAATAGTTGTGAAATCCTCGACTATCACATCCCAAAGGTGCAACTC : 597

TrF3'Ha1 : ----- : -
TrF3'Ha2 : TCTTGG : 603

```

FIGURE 82

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TrPALa : GNAGGAAATTTCAACTAAATATTGCCTTTAATTCTTTNTNATANATNTTTGAATTCNTT : 60  
 TrPALa : CTCCCTAAAAATTCTATAGCTACCACATCANCACAACATAACANNAATTAAGAAATATTN : 120  
 TrPALa : TATNTACTATTTTAAGATATGGAAGTAGTAGCAGCAGCAATCACAAAAACAATGGCAAG : 180  
 TrPALa : ATTGATTCATTTTGCTTGAATCATGCTAATGCTAATAACATGAAAGTGAATGGTGCTGAT : 240  
 TrPALa : CCTTTGAATTGGGGTGTGGCTGCTGAGGCAATGAAGGGAAGTCACTTGGATGAGGTGAAG : 300  
 TrPALa : CGTATGGTGGAGGAATACCGGAAACCGGTTGTCCGTCTTGGTGGCGAGACACTGACGATT : 360  
 TrPALa : TCTCAGGTGGCTGCCATTGCTGCACACGATGGTGCAACGGTGGAGCTATCGGAATCTGCT : 420  
 TrPALa : AGAGCCGGCGTTAAGGCGAGCAGTGACTGGGTTATGGAGAGTATGAACAAAGGTACAGAC : 480  
 TrPALa : AGTTATGGTGTCACTACAGGGTTCGGCGCTACCTCGCACCGCCGAACCAACAAGGTGGT : 540  
 TrPALa : GCTTTGCAGAAAGAGCTCATAAGGTNTTTTGAATGCAGGAATATTTGGAAATGGAACNTG : 600  
 TrPALa : AGACAAAGCCACACACTACCC : 621

## FIGURE 83

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TrPALa : MEVVA<sup>\*</sup>AAITKNNGKIDS<sup>20</sup>FC<sup>\*</sup>LNHANAN<sup>40</sup>NMKVNGADPLNWGVAAEAMKGS<sup>\*</sup>HLDEVKRMVEEY<sup>60</sup> : 60

TrPALa : RKPVVRLGGETLTISQVAAIAAHDGATVELSESARAGVKASSDWVMESMNKG<sup>\*</sup>TDSYGVT<sup>100</sup>T : 120

TrPALa : GFGATSHRRTKQGGALQKELIRFECRNIWKWNRQSHTLP<sup>\*</sup> : 159

## FIGURE 84

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```

      *           20           *           40           *           60
TrPALa1 : GNNGGAAATTNCAACTAAATATTGCCTTTAATTCTTTNTNATANATNTTTGAATTTCCCTT : 60
TrPALa2 : GNAGGAAATTACAACTAAATATTNCCTTTAATTCTTTATNATANATNTTTGAATTTCCNTT : 60
TrPALa3 : -----TCAAGAAATTACACCTCTTTNNTTCTTTNTAATNTTTCCTTTNATTTCNTT : 50

      *           80           *           100          *           120
TrPALa1 : CTCCCTAAAAATTCTATAGCTACCACATCANCACAACATAACANNAATTAAGAAATATTN :120
TrPALa2 : CTCCCTAAAAATTCTATAGCTACCACATCANCACAACATAACANNAATTAAGAAATATTN :120
TrPALa3 : CTCCTCTNGAAATCTATAGCTACCAAAACATAACAAAGTAACACTTATTACTAGGTATTA :110

      *           140          *           160          *           180
TrPALa1 : TATNTACTATTTTAAGATATGGAAGTAGTAGCANAGCAATCACAAAAACAATGGCAAG :180
TrPALa2 : TATNTACTATTTTAAGATATGGAAGTAGTAGCAGCAGCAATCACAAAAACAATGGCAAG :180
TrPALa3 : TATNTACTATTTTAAGATATGGAAGTAGTAGCAGCAGCAATCACAAAAACAATGGCAAG :170

      *           200          *           220          *           240
TrPALa1 : ATTGATTCAATTTTGCTTGAATCATGCTAATGCTAATAACATGAAAGTGAATGGTGCTGAT :240
TrPALa2 : ATTGATTCAATTTTGCTTGAATCATGCTAATGCTAATAACATGAAAGTGAATGGTGCTGAT :240
TrPALa3 : ATTGATTCAATTTTGCTTGAATCATGCTAATGCTAATAACATGAAAGTGAATGGTGCTGAT :230

      *           260          *           280          *           300
TrPALa1 : CCTTTGAATTGGGGTGTGGCTGCTGAGGCAATGAAGGGAAGTCACTTGGATGAGGTGAAG :300
TrPALa2 : CCTTTGAATTGGGGTGTGGCTGCTGAGGCAATGAAGGGAAGTCACTTGGATGAGGTGAAG :300
TrPALa3 : CCTTTGAATTGGGGTGTGGCTGCTGAGGCAATGAAGGGAAGTCACTTGGATGAGGTGAAG :290

      *           320          *           340          *           360
TrPALa1 : CGTATGGTGGAGGAATACCGGAAACCGTTGTCCGTCTTGGTGGCGAGACACTGACGATT :360
TrPALa2 : CGTATGGTGGAGGAATACCGGAAACCGTTGTCCGTCTTGGTGGCGAGACACTGACGATT :360
TrPALa3 : CGTATGGTGGAGGAATACCGGAAACCGTTGTCCGTCTTGGTGGCGAGACACTGACGATT :350

      *           380          *           400          *           420
TrPALa1 : TCTCAGGTGGCTGCCATTGCTGCACACGATGGTGCAACGGTGGAGCTATCGGAATCTGCT :420
TrPALa2 : TCTCAGGTGGCTGCCATTGCTGCACACGATGGTGCAACGGTGGAGCTATCGGAATCTGCT :420
TrPALa3 : TCTCAGGTGGCTGCCATTGCTGCACACGATGGTGCAACGGTGGAGCTATCGGAATCTGCT :410

      *           440          *           460          *           480
TrPALa1 : AGAGCCGGCGTTAAGGCGAGCAGTGACTGGGTTATGGAGAGATGAACAAAGGTACAGAC :480
TrPALa2 : AGAGCCGGCGTTAAGGCGAGCAGTGACTGGGTTATGGAGAGATGAACAAAGGTACAGAC :480
TrPALa3 : AGAGCCGGCGTTAAGGCGAGCAGTGACTGGGTTATGGAGAGATGAACAAAGGTACAGAC :470

      *           500          *           520          *           540
TrPALa1 : AGTTATGGTGTCACTACAGGGTTCCGGCGCTACCTCCACCCGCCGAACCAAACAAGGTGGT :540
TrPALa2 : AGTTATGGTGTCACTACAGGGTTCCGGCGCTACCTCCACCCGCCGAACCAAACAAGGTGGT :540
TrPALa3 : AGTTATGGTGTCACTACAGGGTTCCGGCGCTACCTCCACCCGCCGAACCAAACAAGGTGGT :530

      *           560          *           580          *           600
TrPALa1 : GCTTTGCAGAAAGAGCTCATAAGGTATTTTGTGTTGTCGAAT----- :582
TrPALa2 : GCTTTGCAGAAAGAGCTCATAAGGTNTTTTGAATGCTTGAATATTTGGAAATGGAACNTG :600
TrPALa3 : GCTTTGCAGAAAGAGCTCATAAGGTNTTTTGAATGCAGGAATATTTGGAAATGGAACNTG :590

      *           620
TrPALa1 : ----- : -
TrPALa2 : AGTCNAAGCCACACACTACCC :621
TrPALa3 : ANACAAATCC----- :600

```

FIGURE 85

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TrPALb : GNAGGAAANAATTNTATTGTTATTATTTCCCCCACACAACGGAAANAATTNTATTGTTN : 60

TrPALb : CTTATTTCCCCCACACAACATAACNAATACATTNTCCTCTCCTCTCATCACAAATTATTA : 120

TrPALb : CTTTCTACACACCCCCCTCTCAACTATTATTAACATAATGGAGGGAATTACCAATG : 180

TrPALb : GCCATGCTGAAGCAACTTTTTGCGTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG : 240

TrPALb : CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAAT : 300

TrPALb : ACCGTAATCCATTGGTTAAATTTGGCGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA : 360

TrPALb : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAGGGCCGGCGTTA : 420

TrPALb : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA : 480

TrPALb : CCACCGGTTTCGGCGCCACCTCTCACCGGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG : 540

TrPALb : AGCTAATTAGGTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC : 600

TrPALb : TACCACACACAGCAACCAGAGCTGCAATGCTTGTGAGAATCAACACTCTTCTTCAAGAGG : 660

TrPALb : AATATTTCTTGAATGGCCTTTGTAAATTTTGG : 693

## FIGURE 86

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TrPALb : MEGITNGHAEATFCVTKSVGDPLNWGAAAESLMGSHLDEVKRMVEEYRNPLVKIGGETLT : 60

TrPALb : IAQVAGIASHDSGVRVELSESARAGVKASSDWVMDSMNNGTDSYGVTTGFGATSHRRTKQ : 120

TrPALb : GGALQKELIRFLNAGIFGNGTESNCTLPHTATRAAMLVRINTLLQEYFNLGLCKFL : 177

## FIGURE 87



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		*	20	*	40	*	60	
TrPALb1 :	GNAGGAAANAATTNTATTGTTATTATTTCCCCCACACAAC							: 60
TrPALb2 :	-----							: 21
TrPALb3 :	-----							: 21
TrPALb4 :	-----							: 19
TrPALb5 :	-----							: 19
TrPALb6 :	-----							: 18
TrPALb7 :	-----							: 18
TrPALb8 :	-----							: -
		*	80	*	100	*	120	
TrPALb1 :	CTTTCTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 120
TrPALb2 :	TTT TTTCTACCCACACAACATAACAAATACATTNTCTCTCTCTCATCACAATTATTA							: 81
TrPALb3 :	CCTCTCTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 81
TrPALb4 :	NTT TTTCTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 79
TrPALb5 :	CTTATTTTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 79
TrPALb6 :	ATTATTTTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 78
TrPALb7 :	CTTATTTTCTCTCACANATATACNNTTTCNTAGNACNCTNNCTCTCAACTATTA							: 78
TrPALb8 :	-----							: 42
		*	140	*	160	*	180	
TrPALb1 :	TTTCTAC							: 180
TrPALb2 :	CTTTCTAC							: 141
TrPALb3 :	CTTTCTAC							: 141
TrPALb4 :	CTTTCTAC							: 139
TrPALb5 :	CTTTCTAC							: 139
TrPALb6 :	CTTTCTAC							: 138
TrPALb7 :	CTTTCTAC							: 138
TrPALb8 :	CTTTCTAC							: 102
		*	200	*	220	*	240	
TrPALb1 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 240
TrPALb2 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 201
TrPALb3 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 201
TrPALb4 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 199
TrPALb5 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 199
TrPALb6 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 198
TrPALb7 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 198
TrPALb8 :	GCCATGCTGAAGCAACTTTTTCGCTGACCAAAAGTGTTGGTGATCCACTCAACTGGGGTG							: 162
		*	260	*	280	*	300	
TrPALb1 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 300
TrPALb2 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 261
TrPALb3 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 261
TrPALb4 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 259
TrPALb5 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 259
TrPALb6 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 258
TrPALb7 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 258
TrPALb8 :	CAGCCGCGGAGTCGTTGATGGGGAGTCATTTGGATGAGGTGAAGCGTATGGTGGAGGAGT							: 222
		*	320	*	340	*	360	
TrPALb1 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 360
TrPALb2 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 321
TrPALb3 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 321
TrPALb4 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 319
TrPALb5 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 319
TrPALb6 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 318
TrPALb7 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 318
TrPALb8 :	ACCGTAATCCATTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTCAGGTGGCTGGAA							: 282

FIGURE 88

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      *           380           *           400           *           420
TrPALb1 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :420
TrPALb2 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :381
TrPALb3 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :381
TrPALb4 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :379
TrPALb5 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :379
TrPALb6 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGNNCGAGTCCGCCAAGGGCCGGCGTTA :378
TrPALb7 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :378
TrPALb8 : TTGCTTCTCATGATAGTGGTGTGAGGGTGGAGCTGTCTGAGTCCGCCAAGGGCCGGCGTTA :342

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      *           440           *           460           *           480
TrPALb1 : AGGCGAGTAGTGATTGGGTGATGGAGCATGAACAATGGGACTGATAGTTACGGTGTTA :480
TrPALb2 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :441
TrPALb3 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :441
TrPALb4 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :439
TrPALb5 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :439
TrPALb6 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :438
TrPALb7 : AGGCGAGTAGTGATTGGGTGATGGACAGCATGAACAATGGGACTGATAGTTATGGTGTTA :438
TrPALb8 : AGGCGAGTAGTGATTGGGTGATGGAGCATGAACAATGGGACTGATAGTTATGGTGTTA :402

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```

      *           500           *           520           *           540
TrPALb1 : CCACCGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :540
TrPALb2 : CCACCGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :501
TrPALb3 : CCACCGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :501
TrPALb4 : CCACCGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :499
TrPALb5 : CCACGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :499
TrPALb6 : C----- :441
TrPALb7 : CCACGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :498
TrPALb8 : CCACCGGTTTGGGCGCCACCTCTCACC GGAGAACCAAGCAGGGTGGTGCCTTGCAGAAGG :462

```

```

      *           560           *           580           *           600
TrPALb1 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAN----- :592
TrPALb2 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :561
TrPALb3 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :561
TrPALb4 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :559
TrPALb5 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :559
TrPALb6 : ----- : -
TrPALb7 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :558
TrPALb8 : AGCTAATTAGGTTTTTTGAATGCTGGAATATTTGGCAATGGTACAGAATCTAACTGTACAC :522

```

```

      *           620           *           640           *           660
TrPALb1 : ----- : -
TrPALb2 : TACCACACACAGCAACCAGAGCTGCAATGCTTGTGAGAATCAACACTCTTCTTCAAG--- :618
TrPALb3 : TCCMACACACAGCAACCAGAGCTGCAATGCTTGTGAGAATCAACACTCTTCTTCAAG--- :621
TrPALb4 : TACCACACACAGCAACCAGAGCTGCAATGCTTGTGAGAATCAACACTCTTCTTCAAG--- :590
TrPALb5 : TACCACACACAGCAACCAGAGCTGCAATGCTTGTGAGAATCAACACTCTTCTTCAAG--- :616
TrPALb6 : ----- : -
TrPALb7 : TACCACACC----- :567
TrPALb8 : TACCACACAC----- :532

```

```

      *           680           *
TrPALb1 : ----- : -
TrPALb2 : ----- : -
TrPALb3 : AATATTTCTTGAATGGCCTTTGTAAATTTTTTGG :654
TrPALb4 : ----- : -
TrPALb5 : ----- : -
TrPALb6 : ----- : -
TrPALb7 : ----- : -
TrPALb8 : ----- : -

```

FIGURE 88 (cont)

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TrPALc : AACAAGATCGTTATGCCTTAGAACTTCACCTCAATGGCTTGGTCCTTTGATTGAAGTGAT : 60

TrPALc : AAGATTTTCAACCAAATCAATTGAAAGAGAAATTAACCTCGGTCAACGACAACCCTTTGAT : 120

TrPALc : CGATGTTTCAAGGAACAAGGCCATTCATGGTGGTAACTTTCAAGGAACACCTATTGGAGT : 180

TrPALc : TTCAATGGATAACACACGTTTAGCTCTTGCTTCAATTGGTAAACTCATGTTTGCTCAATT : 240

TrPALc : CTCTGAAC TTGTTAATGATTTTACAACAACGGGTTGCCTTCGAATCTTACTGCTAGTAG : 300

TrPALc : GAACCCGAGCTTGGACTATGGTTTCAAGGGATCGGAAATTGCCATGGCTTCGTATTGTTTC : 360

TrPALc : CGAGTTACAATATCTTGCTAATCCTGTCACCACCCATGTCCAAAGTGCCGAGCAACACAA : 420

TrPALc : CCAAGATGTTAACTCTTTGGGTTTGATTTCATCTAGAAAAACAAATGAAGCTATTGAGAT : 480

TrPALc : TCTCAAGCTCATGTCTTCCACTTTCTTGATTGCATTATGTCAAGCAATCGACTTAAGGCA : 540

TrPALc : CTTGGAGGAAAATCTCAGGAACACCGTCAAGAACACGGT : 579

## FIGURE 89

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TrPALc :                   \*                  20                  \*                  40                  \*                  60  
                  TRSLCLRTSPQWLGPLIEVIRFSTKSIEREINSVNDNPLIDVSRNKAIHGGNFQGTPIGV :   60

TrPALc :                   \*                  80                  \*                  100                  \*                  120  
                  SMDNTRLALASIGKLMFAQFSELVNDFYNNGLPSNLTASRNPSLDYGFKGSEIAMASYCS : 120

TrPALc :                   \*                  140                  \*                  160                  \*                  180  
                  ELQYLANPVTTHVQSAEQHNQDVNSLGLISSRKTNEAIEILKLMSSSTFLIALCQAIDLRH : 180

TrPALc :                   \*  
                  LEENLRNTVKNT : 192

## FIGURE 90

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TrPALd : GGTCAATNCAGCTTNGGAGATCTAGTCCCCCTTCTTACTNTGCTGGTTTACTAACTGGA : 60  
 TrPALd : AGACCNAATTCTAAAGCTCATGGGCCTACAGGAGAAGTACTTAATGCAAAAGAAGCTTTT : 120  
 TrPALd : CAATTGGCTGGAATCAATACCGAGTTCTTTGAATTACAACCAAAAGAAGGTCTTGCACTT : 180  
 TrPALd : GTTAATGGAAGCTGCTGTTGGTTCTGGTTTAGCTTCTATTGTTCTTTTGGAGGCTAACATA : 240  
 TrPALd : TTGGCGGTGTTGTCTGAAGTTCTATCGGCAATTTTCGCTGAAGTTATGCAAGGGAAGCCC : 300  
 TrPALd : GAATTTACTGATCATTTGACACATAAGTTGAAGCACCAACCTGGTCAAATTGAGGCTGCT : 360  
 TrPALd : GCTATTATGGAACACATTTTGGATGGGAGTGCTTATGTTAAAGACGCGAAGAAGTTGCAT : 420  
 TrPALd : GAGATGGACCCTTTACAGAAGCCAAAGCAAGATAGATATGCACTTAGAACTTCACCACAA : 480  
 TrPALd : TGGCTTGGTCCTTTGATTGAAGTGATTAGATTTTCAACCAAGTCAATTGAGAGAGAGATC : 540  
 TrPALd : AACTCTGTCAATGACAACCCCTTTGATTGATGTTTCGAGAAACAAGGCTTTG : 591

## FIGURE 91

# 160/271

TrPALd : GQXSXGDLVPLSYXAGLLTGRXNSKAHGPTGEVLNAKEAFQLAGINTEFFELQPKEGLAL : 60

TrPALd : VNGTAVGSGGLASIVLFEANILAVLSEVLSAIFAEVMQKPEFTDHLTHKLKHHPGQIEAA : 120

TrPALd : AIMEHILDGSAYVKDAKKLHEMDPLQKPKQDRYALRTSPQWLGPLIEVIRFSTKSIEREI : 180

TrPALd : NSVNDNPLIDVSRNKAL : 197

## FIGURE 92

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```

      *           20           *           40           *           60
TrPALe : GNNGGAAATTNCAACTCNATTTTCTTTTNTATAATNTTTGAATTTTCCTTCTCTCTCAAA : 60

      *           80           *           100          *           120
TrPALe : TTCTATAGCTACTCTACCACATCACACAACATAACAAATTAAGAAATATTCATTACTATA : 120

      *           140          *           160          *           180
TrPALe : CTATTAAGATATGGAAGTAGTAGCAGCAGCAATCACAAAAACAACGGCAAGATTGATTG : 180

      *           200          *           220          *           240
TrPALe : ATTTTGCTTGAATCATGCTAATGCTAATAACATGAAAGTGAATGATGCTGATCCTTTGAA : 240

      *           260          *           280          *           300
TrPALe : TTGGGGTGTGGCTGCTGAGGCAATGAAGGGAAGTCACTTGAGATGAGGTGAAACGTATGGT : 300

      *           320          *           340          *           360
TrPALe : GGAGGAGTACCGGAAGCCGATTGTCCGTCTTGGTGGCGAGACGCTGACGATTTCTCAGGT : 360

      *           380          *           400          *           420
TrPALe : GGCTGCCATTGCTGCACACGATGGTGCGATGGTTGAGCTGTCGGAATCTGCTAGAGCCGG : 420

      *           440          *           460          *           480
TrPALe : CGTTAAGGCAAGCAGTGATTGGGTTATGGAGAGTATGAACAAAGGTACTGACAGTTATGG : 480

      *           500          *           520          *           540
TrPALe : TGTCACCACAGGGTTCGGCGCTACCTCNCACCGCCGAACCAACAAGGTGGTGCTTTACA : 540

      *           560          *           580          *
TrPALe : GAAAGGGCTCATAAGGTTTTTGAATGCTGGAATATTTGNAAATGNAACTGAN : 592

```

## FIGURE 93

# 162/271

TrPALe : MEVVAAAITKNNGKIDSFCLNHANANNMKVNDADPLNWGVAAEAMKGSHLDEVKRMVEEY : 60

TrPALe : RKPIVRLGGETLTISQVAAIAAHDGAMVELSESARAGVKASSDWMESMNKGTDSYGVT : 120

TrPALe : GFGATXHRRTKQGGALQKGLIRFLNAGIFXNXTX : 154

## FIGURE 94



# 163/271

```

      *           20           *           40           *           60
TrPALf : CNATTGTTAGTNGTTTCCNCCCACCCACATAACAAATACATAATTCTCTCCTCTGATCAC : 60

      *           80           *           100          *           120
TrPALf : AATTATTACTTTACTACACCCTCCTCTCAACTATTATTAAC TAGCATAATGGAGGGAATT : 120

      *           140          *           160          *           180
TrPALf : ACCAATGGCCATGCTGAAACAAC TTTTAGCGTGACCAAAGTGNNGGNGATCCACTCAAC : 180

      *           200          *           220          *           240
TrPALf : TGGCGNGCAGCCGCGGAGTCGTCGACGGGGAGTCATTTGGATGAGGTGAAGCGTATGGNG : 240

      *           260          *           280          *           300
TrPALf : GAGGAGTACCGTAATCCGNTGGTTAAAATTGGCGGCGAGACGCTTACCATTGCTNNGGTA : 300

      *           320          *           340          *           360
TrPALf : NCTGGAATTGCTTCTCATGATAGTGGAGTGAGGGTGGAGCTGTCCGAGTTCGCAAGGGCC : 360

      *           380          *           400          *           420
TrPALf : GCGGTTAAGGCGAGTAGTGATTGNGTGATGGATAGCATGAACAATGGGACTGATAGTTAC : 420

      *           440          *           460          *           480
TrPALf : GGTGTTACCACCGCNTTTGGTGCCACCTGTCACCGGAGAACCAAGCCANGGTGGTGCCTT : 480

      *           500          *           520          *           540
TrPALf : GCAGAAGGAGCTAAATTNNGGTGTTTTGAANGCTGGNAATANTTTGGCNNTGGTTCAGAA : 540

      *           560
TrPALf : ATCTNAACTTG TNCACTTACCACACC : 566

```

## FIGURE 95

# 164/271

TrPALf : MEGITNGHAETTFSVTKSXXDPLNWXAAAESSTGSHLDEVKRMXEEYRNPXVKIGGETLT : 60

TrPALf : IAXVXGIAHDSGVRVELSEFARAGVKASSDXVMDSMNNGTDSYGVTTXFGATCHRRTKP : 120

TrPALf : XWCLAEGAKXXCFEXWXXFGXGSEIXTCXLTT : 152

## FIGURE 96

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```

      *           20           *           40           *           60
TrPALf1 : CNATTGTTAGTNGTTTCCNCCCACCNACATAACNAATACNTANTTCTCTCCTCTGATCAC : 60
TrPALf2 : -----CCATAACAAATACATTATTCTCTCCTCTGATCAC : 35

      *           80           *           100          *           120
TrPALf1 : AATTATTACTTTTCTACACCCTCCTCTCAACTATTATTAAGTAGCATAATGCAGGGAATT :120
TrPALf2 : AATTATTACTTTTCTACACCCTCCTCTCAACTATTATTAAGTAGCATAATGCAGGGAATT : 95

      *           140          *           160          *           180
TrPALf1 : ACCAATGGCCATGCTGAAACAACCTTTTTCGTGACCAAAAGTGTCGCGGATGTTTCTGTTT :180
TrPALf2 : ACCAATGGCCATGCTGAAACAACCTTTTTCGTGACCAAAAGTGTCGCGGATGTTTCTGTTT :155

      *           200          *           220          *           240
TrPALf1 : TGTCGNG----- :188
TrPALf2 : TGTCGTCGAGCCCGGAGTCGTCGACGGGGAGTCATTTGGATGAGGTGAAGCGTATGGNG :215

      *           260          *           280          *           300
TrPALf1 : ----- : -
TrPALf2 : GAGGAGTACCGTAATCCGNTGGTTAAATTTGGCGGCGAGACGCTTACCATTGCTNNGGTA :275

      *           320          *           340          *           360
TrPALf1 : ----- : -
TrPALf2 : NCTGGAATTGCTTCTCATGATAGTGGAGTGAGGGTGGAGCTGTCCGAGTTCCGAAGGGCC :335

      *           380          *           400          *           420
TrPALf1 : ----- : -
TrPALf2 : GGCGTTAAGGCGAGTAGTGATTGNGTGATGGATAGCATGAACAATGGGACTGATAGTTAC :395

      *           440          *           460          *           480
TrPALf1 : ----- : -
TrPALf2 : GGTGTTACCACCGCCTTTGGTGCCACCTGTCACCGGAGAACCAAGCCANGGTGGTGCCTT :455

      *           500          *           520          *           540
TrPALf1 : ----- : -
TrPALf2 : GCAGAAGGAGCTAAATTNNGGTGTTTGAANGCTGGNAATANTTTGGCNNTGGTTCAGAA :515

      *           560
TrPALf1 : ----- : -
TrPALf2 : ATCTNAACTTGTNCACTTACCACACC :541

```

## FIGURE 97

# 166/271

```

      *           20           *           40           *           60
TrVRa : GTAAGAGTTGAGAAAAANACCAATAAAGTAAACNCTATNTAGAAAGAGAGTCAAAAATG : 60

      *           80           *           100          *           120
TrVRa : GCTGAAGGAAAAGGAAGGGTTTGTGTTACTGGAGGAACAGGTTTCTTGTTTCATGGATC : 120

      *           140          *           160          *           180
TrVRa : ATCAAGAGTCTTCTTGAAAATGGATACTCTGTTAATACCACTATTAGAGCTGATCCAGAA : 180

      *           200          *           220          *           240
TrVRa : CGTAAGAGGGATGTAAGCTTCCTAACAAATCTACCCGGCGCATCCGAAAGGCTACATTTTC : 240

      *           260          *           280          *           300
TrVRa : TTCAACGCCGATCTAGACGACCCAGAGAGTTTCAACGAAGCAATTGAAGGTTGTGTCGGG : 300

      *           320          *           340          *           360
TrVRa : ATATTCCACACCGCTTCACCAATCGATTTGCGCGTGAGTGAGCCAGAAGAAATAGTGACA : 360

      *           380          *           400          *           420
TrVRa : AAAAGAACAGTGGATGGAGCATTAGGAATTTTAAAGCATGTGTGAATTCAAAGACAGTG : 420

      *           440          *           460          *           480
TrVRa : AAGAGATTTATTTACACTTCAAGNGGTTCTGCTGTTTCATTCAATGGAAAAACAAAGAT : 480

      *           500          *           520          *           540
TrVRa : GNNTNGGATGAGAGTGATTGGAGTGATGTTGATTGCTTAGAAGTGTTAAACCATTGTTGGT : 540

      *           560          *           580          *           600
TrVRa : TGGAGTTATGGNGTGTTCAAGACTTTGGCTGAGAAAGCAGTGCTTGAATTTGGTCNACAA : 600

      *           620          *           640          *           660
TrVRa : AATGGGATTGATGTTGTTACTTTGATTCTTCCTTTTATTGTTGGAGGTTTTGTTTGTCCC : 660

      *           680          *           700          *           720
TrVRa : AAGCTTCCTGATTCTGTTGAGAAAGCTCTTGTTTTGGTACTAGGCAAAAAGGAACAAATT : 720

      *           740          *           760          *           780
TrVRa : GGTATTATAAGTTTCCACATGGTACATGTAGATGATGTGGCTAGAGCACATATCTATCTA : 780

      *           800          *           820          *           840
TrVRa : CTTGAGAATCCTGTTCCAGGAGGTAGATATAATTGTTCCACCATTCTTTGTATCTATTGAA : 840

      *           860          *           880          *           900
TrVRa : GAAATGTCACAGCTTCTCTCAGCCAAATATCCAGAATATCAAATACTATCAGTAGATGAG : 900

      *           920          *           940          *           960
TrVRa : TTGAAGGAAATTAAAGGGGCAAGATTGCCAGATTTGAACTCGAAGAAGCTCGTGGACGCT : 960

      *           980          *           1000         *           1020
TrVRa : GGTTTTGAGTTTAAGTATAGTGTCGATGATATGTTGATGATGCGATTCAATGCTGCAAG : 1020

      *           1040         *           1060         *           1080
TrVRa : GAAAAAGGCTATCTCTAAGCATGTATTTGAAAATCCATGAAGTTGAGAAAACAATAATG : 1080

      *           1100         *           1120         *           1140
TrVRa : TGCCTAAAATCAATGATGGCTAATGAGATGTACAAGTTTATGCATTAAGTTATTTGTGAT : 1140

      *           1160         *           1180
TrVRa : CAATCAAATAATGAAATAATCTGTTTCATTTTCCGAAAAAAAAA : 1185

```

## FIGURE 98

# 167/271

TrVRa : MAEGKGRVCVTGGTGFLGSWIIKSLLENGYSVNTTIRADPERKRDVSFLTNLPGASERLH : 60

TrVRa : FFNADLDDPESFNEAIEGCVGIFHTASPIDFAVSEPEEIVTKRTVDGALGILKACVNSKT : 120

TrVRa : VKRFIYTSXGSAVSFNGKNKDXXDESDWSDVDLLRSVKPFGWSYXVFKTLAEKAVLEFGX : 180

TrVRa : QNGIDVVTLLLPFIVGGFVCPKLPDSVEKALVLVLGKKEQIGIISFHMVHVDDVARAHY : 240

TrVRa : LLENPVPGGRYNCSPFFVSIEEMSQLLSAKYPEYQILSVDELKEIKGARLPDLNSKKLVD : 300

TrVRa : AGFEFKYSVDDMFDDAIQCCKEKGYL : 326

## FIGURE 99

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		*	20	*	40	*	60	
TrVRa1 :	<b>GTA</b>	<b>TTAGTTGAGAAAAAA</b>	<b>TAC</b>	<b>NAATAAAAGTAAACNCTATNTAGAAAAGAGAGTCAAAAATG</b>	:	60		
TrVRa2 :	---	<b>AGAGTTGAGAAAAAA</b>	<b>N</b>	<b>CCAATAAAAGTAAACNCTATNTAGAAAAGAGAGT</b>	:	57		
TrVRa3 :	-----				:	-		
TrVRa4 :	-----				:	-		
TrVRa5 :	-----				:	-		
TrVRa6 :	-----				:	-		

		*	80	*	100	*	120	
TrVRa1 :	<b>GCTGAAGGAAAAGGAAGGGTTTGTGTTACTGGAGGAACAGGTTTTCTTGGGTTCATGGATC</b>	:	120					
TrVRa2 :	<b>GCTGAAGGAAAAGGAAGGGTTTGTGTTACTGGAGGAACAGGTTTTCTTGGGTTCATGGATC</b>	:	117					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

		*	140	*	160	*	180	
TrVRa1 :	<b>ATCAAGAGTCTTCTTGAAAATGGATACTCTGTTAATACCACTATTAGAGCTGATCCAGAA</b>	:	180					
TrVRa2 :	<b>ATCAAGAGTCTTCTTGAAAATGGATACTCTGTTAATACCACTATTAGAGCTGATCCAGAA</b>	:	177					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

		*	200	*	220	*	240	
TrVRa1 :	<b>CGTAAGAGGGATGTAAGCTTCCTAACAATCTACCCGGCGCATCCGAAAGGCTACATTTTC</b>	:	240					
TrVRa2 :	<b>CGTAAGAGGGATGTAAGCTTCCTAACAATCTACCCGGCGCATCCGAAAGGCTACATTTTC</b>	:	237					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

		*	260	*	280	*	300	
TrVRa1 :	<b>TTCAACGCCGATCTAGACGACCCAGAGAGTTTCAACGAAGCAATTGAAGGTTGTGTCTGGG</b>	:	300					
TrVRa2 :	<b>TTCAACGCCGATCTAGACGACCCAGAGAGTTTCAACGAAGCAATTGAAGGTTGTGTCTGGG</b>	:	297					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

		*	320	*	340	*	360	
TrVRa1 :	<b>ATATTCCACACCGCTTCACCAATCGATTTCGCCGTGAGTGAGCCAGAAGAAATAGTGACA</b>	:	360					
TrVRa2 :	<b>ATATTCCACACCGCTTCACCAATCGATTTCGCCGTGAGTGAGCCAGAAGAAATAGTGACA</b>	:	357					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

		*	380	*	400	*	420	
TrVRa1 :	<b>AAAAGAACAGTGGATGGAGCATTAGGAATTTTAAAAGCATGTGTGAATTCAAAGACAGTG</b>	:	420					
TrVRa2 :	<b>AAAAGAACAGTGGATGGAGCATTAGGAATTTTAAAAGCATGTGTGAATTCAAAGACAGTG</b>	:	417					
TrVRa3 :	-----	:	-					
TrVRa4 :	-----	:	-					
TrVRa5 :	-----	:	-					
TrVRa6 :	-----	:	-					

FIGURE 100

		*	440	*	460	*	480	
TrVRa1 :	AAGAGATTTATTTACACTTCAAGNGGTTCTGCTGTTTCATTCAATG	AAAAANCAAAGAT	:	480				
TrVRa2 :	AAGAGATTTATTTACACTTCAAGTGGTTCTGCTGTTTCATTCAATG	AAAAAACAAGAT	:	477				
TrVRa3 :	-----	-----	:	-				
TrVRa4 :	-----	-----	:	-				
TrVRa5 :	-----	-----	:	-				
TrVRa6 :	-----	-----	:	-				
		*	500	*	520	*	540	
TrVRa1 :	GNNTNNNATGANA	-----	:	493				
TrVRa2 :	GGTTGGATGAGAGTGAATGGAGTGAATGATTGCTTAGAAGTGTTAAACCATTGGT	-----	:	537				
TrVRa3 :	-----	-----	:	-				
TrVRa4 :	-----	-----	:	-				
TrVRa5 :	-----	-----	:	-				
TrVRa6 :	-----	-----	:	-				
		*	560	*	580	*	600	
TrVRa1 :	-----	-----	:	-				
TrVRa2 :	TGGAGTTATGGTGTTCGAAGACTTTGGCTGAGAAAGCAGTGCTTGAATTTGGTC	ACAA	:	597				
TrVRa3 :	-----GNGTGTTGAAGACTTTGCTGAGAAAGCAGTGCTTGAATTTGGTC	ACAA	:	50				
TrVRa4 :	-----AGACGTTNGCTGAGAAAGCGTGCTTGAATTTGGTC	ACAA	:	41				
TrVRa5 :	-----TTGAATTTGGTC	ACAA	:	19				
TrVRa6 :	-----TTGAATTTGGTC	ACAA	:	19				
		*	620	*	640	*	660	
TrVRa1 :	-----	-----	:	-				
TrVRa2 :	AATG	-----	:	601				
TrVRa3 :	AATGGGATTGATGTTGTTACTTTGATTCTTCCTTTTATTGTTGGAGGTTTTGTTTGTCC	TT	:	110				
TrVRa4 :	AATGGGATTGATGTTGTTACTTTGATTCTTCCTTTTATTGTTGGAGGTTTTGTTTGTCC	TT	:	101				
TrVRa5 :	AATGGGATTGATGTTGTTACTTTGATTCTTCCTTTTATTGTTGGAGGTTTTGTTTGTCCC	TT	:	79				
TrVRa6 :	AATGGGATTGATGTTGTTACTTTGATTCTTCCTTTTATTGTTGGAGGTTTTGTTTGTCCC	TT	:	79				
		*	680	*	700	*	720	
TrVRa1 :	-----	-----	:	-				
TrVRa2 :	-----	-----	:	-				
TrVRa3 :	AAGCTTCCTGATTCTGTTGAGAAAGCTCTTGTTTTGGTACTAGGCAAAAAGGAACAAATT	TT	:	170				
TrVRa4 :	AAGCTTCCTGATTCTGTTGAGAAAGCTCTTGTTTTGGTACTAGGCAAAAAGGAACAAATT	TT	:	161				
TrVRa5 :	AAGCTTCCTGATTCTGTTGAGAAAGCTCTTGTTTTGGTACTAGGCAAAAAGGAACAAATT	TT	:	139				
TrVRa6 :	AAGCTTCCTGATTCTGTTGAGAAAGCTCTTGTTTTGGTACTAGGCAAAAAGGAACAAATT	TT	:	139				
		*	740	*	760	*	780	
TrVRa1 :	-----	-----	:	-				
TrVRa2 :	-----	-----	:	-				
TrVRa3 :	GGTATTATAAGTTTCCACATGGTACATGTGGATGATGTGGCTAGAGCACATATCTATCTA	TT	:	230				
TrVRa4 :	GGTATTATAAGTTTCCACATGGTACATGTGGATGATGTGGCTAGAGCACATATCTATCTA	TT	:	221				
TrVRa5 :	GGTATTATAAGTTTCCACATGGTACATGTAGATGATGTGGCTAGAGCACATATCTATCTA	TT	:	199				
TrVRa6 :	GGTATTATAAGTTTCCACATGGTACATGTAGATGATGTGGCTAGAGCACATATCTATCTA	TT	:	199				
		*	800	*	820	*	840	
TrVRa1 :	-----	-----	:	-				
TrVRa2 :	-----	-----	:	-				
TrVRa3 :	CTTGAGAATCCTGTTCCAGGAGGTAGATATAATTGTTCCACCATTCTTTGTATCTATTGAA	TT	:	290				
TrVRa4 :	CTTGAGAATCCTGTTCCAGGAGGTAGATATAATTGTTCCACCATTCTTTGTATCTATTGAA	TT	:	281				
TrVRa5 :	CTTGAGAATCCTGTTCCAGGAGGTAGATATAATTGTTCCACCATTCTTTGTATCTATTGAA	TT	:	259				
TrVRa6 :	CTTGAGAATCCTGTTCCAGGAGGTAGATATAATTGTTCCACCATTCTTTGTATCTATTGAA	TT	:	259				

**FIGURE 100 (cont)**

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		*	860	*	880	*	900	
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	GAAATGTCACAGCTTCTTTCAGCCAAATATCCAGAATATCAAATACTATCTGTAGATGAG							: 350
TrVRa4 :	GAAATGTCACAGCTTCTTTCAGCCAAATATCCAGAATATCAAATACTATCTGTAGATGAG							: 341
TrVRa5 :	GAAATGTCACAGCTTCTCTCAGCCAAATATCCAGAATATCAAATACTATCAGTAGATGAG							: 319
TrVRa6 :	GAAATGTCACAGCTTCTCTCAGCCAAATATCCAGAATATCAAATACTATCAGTAGATGAG							: 319
		*	920	*	940	*	960	
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	TTGAAGGAAATTAAAGGGGCAAGCTTGCCAGATTTGAACTCGAAGAAGCTCGTGGACGCT							: 410
TrVRa4 :	TTGAAGGAAATTAAAGGGGCAAGCTTGCCAGATTTGAACTCGAAGAAGCTCGTGGACGCT							: 401
TrVRa5 :	TTGAAGGAAATTAAAGGCGCAAGATTGCCAGATTTGAACTCGAAGAAGCTCGTGGACGCT							: 379
TrVRa6 :	TTGAAGGAAATTAAAGCGCAAGATTGCCAGATTTGAACTCGAAGAAGCTCGTGGACGCT							: 379
		*	980	*	1000	*	1020	
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	GGTTTTGAGTTTAAAGTATAGTGTCTCGATGATATGTTTCGATGATGCGATTCAATGCTGCAAG							: 470
TrVRa4 :	GGTTTTGAGTTTAAAGTATAGTGTCTCGATGATATGTTTCGATGATGCGATTCAATGCTGCAAG							: 461
TrVRa5 :	GGTTTTGAGTTTAAAGTATAGTGTCTCGATGATATGTTTCGATGATGCGATTCAATGCTGCAAG							: 439
TrVRa6 :	GGTTTTGAGTTTAAAGTATAGTGTCTCGATGATATGTTTCGATGATGCGATTCAATGCTGCAAG							: 439
		*	1040	*	1060	*	1080	
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	GAAAAAGGCTATCTCTAAGCATGTCTTTGAAAATTCCATGAAGTTGAGAAAACAATAATG							: 530
TrVRa4 :	GAAAAAGGCTATCTCTAAGCATGTCTTTGAAAATTCCATGAAGTTGAGAAAACAATAATG							: 521
TrVRa5 :	GAAAAAGGCTATCTCTAAGCATGTATTTGAAAATTCCATGAAGTTGAGAAAACAATAATG							: 499
TrVRa6 :	GAAAAAGGCTATCTCTAAGCATGTATTTGAAAATTCCATGAAGTTGAGAAAACAATAATG							: 499
		*	1100	*	1120	*	1140	
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	TGCCATAAATCAATGATGGCTAATGAGATGTACAAGTTTATGCATTAAGTTATTTGTGAT							: 590
TrVRa4 :	TGCCATAAATCAATGATGGCTAATGAGATGTACAAGTTTATGCATTAAGTTATTTGTGAT							: 581
TrVRa5 :	TGCCATAAATCAATGATGGCTAATGAGATGTACAAGTTTATGCATTAAGTTATTTGTGAT							: 559
TrVRa6 :	TGCCATAAATCAATGATGGCTAATGAGATGTACAAGTTTATGCATTAAGTTATTTGTGAT							: 559
		*	1160	*	1180			
TrVRa1 :	-----							-
TrVRa2 :	-----							-
TrVRa3 :	CAATCAAATAATGAAATAATCTG-----							: 613
TrVRa4 :	CAATCAAATAATGAAATAATC-----							: 602
TrVRa5 :	CAATCAAATAATGAAAN-----							: 575
TrVRa6 :	CAATCAAATAATGAAATAATCTGTTTCATTTTCCGAAAAAAAAA							: 604

FIGURE 100 (cont)



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```

      *           20           *           40           *           60
LpDFRa : GTSYWTTCGAGTTTGAGAGAATGGCTTCCAGGGCAAGGTGTGTGTTACTGGGGCCTCTGG : 60

      *           80           *           100          *           120
LpDFRa : CTTTGTTGCTTCTTGGCTTGTCAAAAGACTACTCGAGTCCGGTTATAATGTTCTAGGGAC : 120

      *           140          *           160          *           180
LpDFRa : AGTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAA : 180

      *           200          *           220          *           240
LpDFRa : GGAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGAAGAAGGGAGCTTCGATGATGCTGT : 240

      *           260          *           280          *           300
LpDFRa : GATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAA : 300

      *           320          *           340          *           360
LpDFRa : GGAAGAAATGCTTGATTCTGCAATTAACGGCACTCTAAACGTGCTGAGATCGTGCAAGAA : 360

      *           380          *           400          *           420
LpDFRa : GAATCCTTTTTCTCAAAGGGTTGTTCTCACGTCATCATCGTCAACCGTGAGGCTGAGGGA : 420

      *           440          *           460          *           480
LpDFRa : TGAAGCTGAATTCCCACCCAACGTGTTGCTGGATGAAACATCATGGAGCTCCGTGGAGTT : 480

      *           500          *           520          *           540
LpDFRa : CTGTGAAAGTATCCAGGTATGGTATGGTGTCGCGAAGATCCTTGCTGAGAAATCAGCTTG : 540

      *           560          *           580          *           600
LpDFRa : GGAGTTCGCCAAGGAGAACAACATCGACCTAGTGGCTGTTCTTCCAACGTTTCGTGATTGG : 600

      *           620          *           640          *           660
LpDFRa : ACCTAATCTCTCGTCTGAATTAGGACCCACTGTTTTAGATGTCCTTGGCTTATTTAAAGG : 660

      *           680          *
LpDFRa : AGAGACAGAGAAGTTCACCATGTTTKGGAAGGATG : 695

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## FIGURE 101

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LpDFRa : VFSSLREWLPGQVCVTGASGFVASWLVKRLLESGYNVLGTVRDPGNQKKVAHLWNLAGAK : 60

LpDFRa : ERLELVKADLLEEGSFDDAVMACEGVFHTASPIITKSDTKEEMLDSAINGTLNVLRSCCK : 120

LpDFRa : NPFLKRVVLTSSSSTVRLRDEAEFPPNVLLDETSWSSVEFCESIQVWYGVAKILAEKSAW : 180

LpDFRa : EFAKENNIDLVAVLPTFVIGPNLSSELGPTVLDVLGLFKGETEKFTMFGKD : 231

**FIGURE 102**

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          *           20           *           40           *           60
LpDFRa1 : -----CTTTGAGAGAATGGCTTCNAGGGCAGGTGTGTGTACTGGGGCCTCTGGC : 50
LpDFRa2 : -----TCGAGTTTGAGAGAATGGCTTCNAGGGCAGGTGTGTGTACTGGGGCCTCTGGC : 54
LpDFRa3 : -----GTTGGCTTCCAGGGCGGTGTGTGTGTACTGGGGCCTCTGGC : 41
LpDFRa4 : --GCATTTCGAGTTTGAGAGAATGGCTTCCAGGGCAGGTGTGTGTACTGGGGCCTCTGGC : 58
LpDFRa5 : GTCTTTTCGAGTTTGAGAGAATGGCTTCNAGGGCAGGTGTGTGTACTGGGGCCTCTGGC : 60

          *           80           *           100          *           120
LpDFRa1 : TTTGTTGCTTCTTGGCTTGTCAAAAGACTACTCGAGTCCGGTTATAATGTTCTAGGGACA : 110
LpDFRa2 : TTTGTTGCTTCTTGGCTTGTCAAAAGACTACTCGAGTCCGGTTATAATGTTCTAGGGACA : 114
LpDFRa3 : TTTGTTGCTTCTTGGCTTGTNAAA-GACTACTCGAGTCCGGTTATAATGTTCTAGGGACA : 100
LpDFRa4 : TTTGTTGCTTCTTGGCTTGTCAAAAGACTTCTCGAGTCCGGTTATAATGTTCTAGGGACA : 118
LpDFRa5 : TTTGTTGCTTCTTGGCTTGTCAAAAGACTTCTCGAGTCCGGTTATAATGTTCTAGGGACA : 120

          *           140          *           160          *           180
LpDFRa1 : GTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAAG : 170
LpDFRa2 : GTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAAG : 174
LpDFRa3 : GTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAAG : 160
LpDFRa4 : GTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAAG : 178
LpDFRa5 : GTCAGAGACCCAGGCAATCAGAAGAAGGTAGCACACCTCTGGAACCTAGCAGGGGCCAAG : 180

          *           200          *           220          *           240
LpDFRa1 : GAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGGGAAGAAGGGAGCTTCGATGATGCTGTG : 230
LpDFRa2 : GAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGGGAAGAAGGGAGCTTCGATGATGCTGTG : 234
LpDFRa3 : GAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGGGAAGAAGGGAGCTTCGATGATGCTGTG : 220
LpDFRa4 : GAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGGGAAGAAGGGAGCTTCGATGATGCTGTG : 238
LpDFRa5 : GAAAGGTTGGAGCTTGTCAAAGCTGACCTCTTGGGAAGAAGGGAGCTTCGATGATGCTGTG : 240

          *           260          *           280          *           300
LpDFRa1 : ATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAAG : 290
LpDFRa2 : ATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAAG : 294
LpDFRa3 : ATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAAG : 280
LpDFRa4 : ATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAAG : 298
LpDFRa5 : ATGGCCTGTGAGGGTGTCTTCCACACTGCATCACCTATCATCACCAAATCTGATACCAAG : 300

          *           320          *           340          *           360
LpDFRa1 : GAAGAAATGCTTGATTCTGCAATTAACGGCACTCTAAACGTGCTGAGATCGTGCAAGAAG : 350
LpDFRa2 : GAAGAAATGCTTGATTCTGCAATTAACGGCACTCTAAACGTGCTGAGATCGTGCAAGAAG : 354
LpDFRa3 : GAAGAAATGCTTGATTCTGCAATTAACGGCACTCTAAACGNGCTGAGATCGNGCAAGAAG : 340
LpDFRa4 : GAAGAAATGCTTGATTCTGCAATTAACGGCACTCTAAACGTGCTGAGATCGTGCAAGAAG : 358
LpDFRa5 : GAAGAAATGCTTGATTCTGCAATTAACGGCNCCTCTAAACGTGCTGNNATCGGGTNAAG : 360

          *           380          *           400          *           420
LpDFRa1 : AATCCTTTTCTCAAAAGGGTTGTTCTCACGTCATCATCGTCAACCGTGAGGCTGAGGGAT : 410
LpDFRa2 : AATCCTTTTCTCAAAAGGGTTGTTCTCACGTCATCATCGTCAACCGTGAGGCTGAGGGAT : 414
LpDFRa3 : AATNCTTTTCTNAAAAGGNTGNTCTCACGTCATCATCGTCAACCGNGANGCTGANGGAT : 400
LpDFRa4 : AATCCTTTTCTCAAAAGGGTTGTTCTCACGTCATCATCGTCAACCGTGAGGCTGAGGGAT : 418
LpDFRa5 : AATAAN----- : 365

          *           440          *           460          *           480
LpDFRa1 : GAAGCTGAATTCCCACCCAACGTGTTGCTGGATGAAACATCATGGAGCTCCGTGGAGTTC : 470
LpDFRa2 : GAAGCTGAATTCCCACCCAACGTGTTGCTGGATGAAACATCATGGAGCTCCGTGGAGTTC : 474
LpDFRa3 : GAANCTGANTTCCCACCCAACGNGN----- : 425
LpDFRa4 : GAAGCTGAATTCCCACCCAACGTGTTGCTGGATGAAACATCATGGAGCTCCGTGGAGTTC : 478
LpDFRa5 : ----- : -

```

FIGURE 103

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		*	500	*	520	*	540	
LpDFRa1 :	TGTGAAAGTATCCAGGTATGGTATGGTGTCTCGCGAAGATCCTTGCTGAGAAATCAGCTTGG							: 530
LpDFRa2 :	TGTGAAAGTATCCAGGTATGGTATGGTGTCTCGCGAAGATCCTTGCTGAGAAATCAGCTTGG							: 534
LpDFRa3 :	-----							: -
LpDFRa4 :	TGTGAAAGTATCCAGGTATGGTACGGTGTCTCGCAAAGATCCTTGCCGAGAAATCAGCCTGG							: 538
LpDFRa5 :	-----							: -
		*	560	*	580	*	600	
LpDFRa1 :	GAGTTCGCCAAGGAGAAACAACATCGACCTAGTGGCTGTTCTTCCAACGTTCCGTGATTGGA							: 590
LpDFRa2 :	GAGTTCGCCAAGGAGAAACAACATCGACCTAGTGGCTGTTCTTCCAACGTTCCGTGATTGGA							: 594
LpDFRa3 :	-----							: -
LpDFRa4 :	GAGTTTGCCAAGGAGAAACAACATCGACCTAGTGGCTGTTCTTCCAACATTCCGTGATTGGA							: 598
LpDFRa5 :	-----							: -
		*	620	*	640	*	660	
LpDFRa1 :	CCTAATCTCTCGTCTGAATTAGGACCCACTGTTTTAGATGTCCTTGGCTTATTTAAAGGA							: 650
LpDFRa2 :	CCTAATCTCTCTCGTCTGAATTAGGACCCACTGTTTTAGATGTCCTTGGCTTATTTAANGGA							: 654
LpDFRa3 :	-----							: -
LpDFRa4 :	CCTAATCTCTCGTCTGAATTAGGACCCACTGTTTTAGATGTCCTTGGCTTATTTAAAGGA							: 658
LpDFRa5 :	-----							: -
		*	680	*				
LpDFRa1 :	GAGACAGAGAAGTTCAC-----							: 667
LpDFRa2 :	GAGACAGAGAAGTTCACCATGTTTGGGAAGGATG							: 688
LpDFRa3 :	-----							: -
LpDFRa4 :	GAGACAGAGAAGTTCACCATGTTTGGGAAGGAN-							: 691
LpDFRa5 :	-----							: -

FIGURE 103 (cont)

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```

      *           20           *           40           *           60
LpDFRb : GTCCTCGCCTACGAGCGCCCCGACGCCCCGCGCCGCTACCTCTGCATCGGGGCCGTGCTG : 60

      *           80           *           100          *           120
LpDFRb : CACCGCGCGCACTTCCTAAAGCTTCTCAAGGACCTCTTCCCGCAGTACTCCTTCACCGCC : 120

      *           140          *           160          *           180
LpDFRb : AAGTGCGAAGACGACGGCAAGCCCATGGCGAAGCCGTACAAGTTCTCCNACCAGAGGCTC : 180

      *           200          *           220          *           240
LpDFRb : AGGGACCTGGGATTAAAAATTCACCTCCGCTGGCGGAAAGTTTGTACGAGACCGTGACGTGC : 240

      *           260          *           280          *           300
LpDFRb : CTGCAAAAAAATGGCCACCTGCCTCTGCCCCTCCCATGGCGCCAAAGCGTGACATACCTA : 300

      *           320          *           340          *           360
LpDFRb : TAATACTACAAAGACACGGCCGGGATCGACAAGCCAAGAAACAGAGGATTCTCCCGAGGT : 360

      *           380          *           400          *           420
LpDFRb : TCACCATGGAATTGTGTATTTACAAAGTTTGAATTCTTATTTTTTTTATTATGAAGAAA : 420

      *           440          *           460          *           480
LpDFRb : TACGAAAACCAATACTGTATACCAGAGGCAAGTGTAACAATGTAAATAGTCGTGTAAAT : 480

      *           500          *           520
LpDFRb : CTTGTTCAAGAATGAATGATAAAGTATTTTTTGCAAAAAAAAAA : 524

```

## FIGURE 104

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                  \*                  20                  \*                  40                  \*                  60  
LpDFRb : VLAYERPDARGRYLCIGAVLHRAHFLKLLKDLFPQYSFTAKCEDDGKPMAPYKFSXQRL : 60

                  \*                  80                  \*                  100  
LpDFRb : RDLGLKFTPLAESLYETVTCLOKNHGLPLPAPMAPKRAYL : 100

## FIGURE 105

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	*                      20                      *                      40                      *                      60	
LpDFRb1 :	<b>GTCTCGCCTACGAGCGCCCCGACGCCCGCGCCGCTACCTCTGCATCGGGGCCGTGCTG</b>	: 60
LpDFRb2 :	-----	: -
LpDFRb3 :	-----	: -
	*                      80                      *                      100                      *                      120	
LpDFRb1 :	<b>CACCGCGCGCACTTCCTAAAGCTTCTCAAGGACCTCTTCCCGCAGTACTCCTTCACCGCC</b>	: 120
LpDFRb2 :	-----	: -
LpDFRb3 :	-----	: -
	*                      140                      *                      160                      *                      180	
LpDFRb1 :	<b>AAGTGCGAAGACGACGGCAAGCCCATGGCGAAGCCGTACAAGTTCTCCNACCAGAGGCTC</b>	: 180
LpDFRb2 :	-----AAGCCGTACAAGTTCTCCNACCAGAGGCTC	: 30
LpDFRb3 :	-----GTTCTCNACCAGAGGCTC	: 19
	*                      200                      *                      220                      *                      240	
LpDFRb1 :	<b>AGGGACCTGGGATTAAAATTCACTCCGCTGGCGGAAAGTTTGTACGAGACCGTGACGTGC</b>	: 240
LpDFRb2 :	<b>AGGGACCTGGGATTAAAATTCACTCCGCTGGCGGAAAGTTTGTACGAGACCGTGACGTGC</b>	: 90
LpDFRb3 :	<b>AGGGACCTGGGATTAAAATTCACTCCGCTGGCGGAAAGTTTGTACGAGACCGTGACGTGC</b>	: 79
	*                      260                      *                      280                      *                      300	
LpDFRb1 :	<b>CTGCAAAAAAATGGCCACCTGCCTCTGCCCGCTCCCCTGGCGCCAAAGCGTGCATACCTA</b>	: 300
LpDFRb2 :	<b>CTGCAAAAAAATGGCCACCTGCCTCTGCCCGCTCCCCTGGCGCCAAAGCGTGCATACCTA</b>	: 150
LpDFRb3 :	<b>CTGCAAAAAAATGGCCACCTGCCTCTGCCCGCTCCCCTGGCGCCAAAGCGTGCATACCTA</b>	: 139
	*                      320                      *                      340                      *                      360	
LpDFRb1 :	<b>TAATAATACAAAGACACGGCCGGGATCGACAAGCCAAGAAACAGAGGATTCTCCCGAGGT</b>	: 360
LpDFRb2 :	<b>TAATACTACAAAGACACGGCCGGGATCGACAAGCCAAGAAACAGAGGATTCTCCCGAGGT</b>	: 210
LpDFRb3 :	<b>TAATACTACAAAGACACGGCCGGGATCGACAAGCCAAGAAACAGAGGATTCTCCCGAGGT</b>	: 199
	*                      380                      *                      400                      *                      420	
LpDFRb1 :	<b>TCACCATGGAATTGTGTATTTTCAAAAGTTTGAATTCTTATTTTTTTTTATTATGAAGAAA</b>	: 420
LpDFRb2 :	<b>TCACCATGGAATTGTGTATTTTCAAAAGTTTGAATTCTTATTTTTTTTTATTATGAAGAAA</b>	: 270
LpDFRb3 :	<b>TCACCATGGAATTGTGTATTTTCAAAAGTTTGAATTCTTATTTTTTTTTATTATGAAGAAA</b>	: 259
	*                      440                      *                      460                      *                      480	
LpDFRb1 :	<b>TACGGAAACCAATACTGTATACCAGAGGCAAGTGTAACAATGTAAATAGTCGTGTAAAT</b>	: 480
LpDFRb2 :	<b>TACGGAAACCAATACTGTATACCAGAGGCAAGTGTAACAATGTAAATAGTCGTGTAAAT</b>	: 330
LpDFRb3 :	<b>TACGGAAACCAATACTGTATACCAGAGGCAAGTGTAACAATGTAAATAGTCGTGTAAAT</b>	: 319
	*                      500                      *                      520	
LpDFRb1 :	<b>CTTGTTCAGAATGAATGATAAAGTATTTTTTGCAAAAAA</b>	: 524
LpDFRb2 :	<b>CTTGTTCAGAATGAATGATAAAGTATTTTTTGCAAAAAA</b>	: 374
LpDFRb3 :	<b>CTTGTTCAGAATGAATGATAAAGTATTTTTTGCAAAAAA</b>	: 363

FIGURE 106

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```

      *           20           *           40           *           60
LpF3Ha : TCTCNAGACACACTGTGTAACCACGGTAGCGAGTGGCAAGACTAGCAGAAAGTACGGACA : 60

      *           80           *           100          *           120
LpF3Ha : TCAGCTAACCATTCTCTCAACTAGATAAGCATGGCTCCGGCGATGTCCAACCCTCTCCTC : 120

      *           140          *           160          *           180
LpF3Ha : AGTGATCGGGTGGCACGCTCCAAGAAAGTCCCATCTAGCCACGTTAGAGCGGTGGGAGAC : 180

      *           200          *           220          *           240
LpF3Ha : CGCCCAGACCTCGCCAATGTTCGACCACGAGTCCGGCGCGGGCATTCGGCTCATCGACCTG : 240

      *           260          *           280          *           300
LpF3Ha : AAGCAGCTCGAAGGTCCAGGGCGCCGCAGGGTCGTCGAGGCCATCGGCTCCGCGTGCAGAG : 300

      *           320          *           340          *           360
LpF3Ha : AACGATGGGTTTTTTCATGGTGACGAATCATGGCATCCCAGAGGCGGTCTGGAGGGGATG : 360

      *           380          *           400          *           420
LpF3Ha : CTGAGCGTGGCGAGGGAGTTCTTCCACCTGCCGGAGTCGGAGCGGCTCAAGTGCTACTCC : 420

      *           440          *           460          *           480
LpF3Ha : GACGACCCCAAGAAGGCGGTCCGGCTGTTCGACGAGCTTCAACGTGCGCACGGAGAAGGTG : 480

      *           500          *           520          *           540
LpF3Ha : AGCAACTGGCGCGACTTCCTCCGGCTGCATTGCTACCCTCTTGAGAGCTTCGTCGACCAG : 540

      *           560          *           580          *           600
LpF3Ha : TGGCCGTCGAACCCGCGCCCTTCAGGCAAGTCGTCGGCACCTACTCGACGGAAGCGAGA : 600

      *           620          *           640          *           660
LpF3Ha : GCGCTGGCGCTGAGGCTCCTGGAGGCGATATCGGAGAGCCTAGGGCTGGAGAGAGGCCAC : 660

      *           680          *           700          *           720
LpF3Ha : ATGGTGAAGGCCATGGGGCGGCACGCGCAGCACATGGCGGTGAACTACTACCCGCCGTGC : 720

      *           740          *           760          *           780
LpF3Ha : CCGCAGCCGGAGCTCACCTACGGTCTGCCAGGGCACACGGACCCCAACGCCCTCACCATC : 780

      *           800          *           820          *           840
LpF3Ha : CTCCTCATGGATCCCCACGTCTCCGGCCTCCAGGTCCTCAGGGACGGCGCCAAGTGGATC : 840

      *           860          *           880          *           900
LpF3Ha : GCCGTCCACCCACGCCCCAACGCCCTGGTCAACCTAGGCGACCAGCTACAGGCGCTG : 900

      *           920          *           940          *           960
LpF3Ha : AGCAACGGCGCGTACAAGAGCGTGTGGCACCAGGGCAGTGGTGAACGCGGAGCAGGAGCGT : 960

      *           980          *           1000          *           1020
LpF3Ha : CTGTCGGTGGCATCTTTCTGTGCCCCGTGCAACAGCGCGGTTATCTGCCCCGCGCCGAGG : 1020

      *           1040          *           1060          *           1080
LpF3Ha : CTCGTCGGCGACGGGGAGGACCCCGTCTACCGGAGCTACACCTACGACGAGTACTACAAG : 1080

      *           1100          *           1120          *           1140
LpF3Ha : AGGTTTTGGAGCAGGAACCTGGATCAGGAGCACTGCCTCGAGCTCTTCAGGAGTCAGCAC : 1140

```

## FIGURE 107



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```

          *           1160           *           1180           *           1200
LpF3Ha : TGATGCTTGAACCTTGAGTTACTAGCTAGCTCTCCTTAACAGTGCAAATCCATGGCCCAA :1200

          *           1220           *           1240           *           1260
LpF3Ha : GAGGGCCCCGATTGCATGGTTACTTATGTTGTTTGAAGTGGTATTGCTTAAGTGCCTAAT :1260

          *           1280           *           1300           *           1320
LpF3Ha : AACATTGCTACATTCTACTNCTATCTTGTCCGTTTAAAATTATAAGATGGCCTAACCTTT :1320

          *           1340           *           1360           *           1380
LpF3Ha : TTCTTAATTGTATGCATNCTGAACATATTTAAGTGTGTGTGTTTCAGACAGTTTAGTCTGC :1380

LpF3Ha : A :1381
```

## FIGURE 107 (cont)

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\*                      20                      \*                      40                      \*                      60  
 LpF3Ha : MSNPLLSDRVARSKKVPSSHVRAVGDRPDLANVDHESGAGIPLIDLKQLEGPGRRRVVEA : 60

\*                      80                      \*                      100                      \*                      120  
 LpF3Ha : IGSACENDGFFMVTNHGIPEAVVEGMLSVAREFFHLPESERLKCYSDDPKKAVRLSTSFN : 120

\*                      140                      \*                      160                      \*                      180  
 LpF3Ha : VRTEKVSNNWRDFLRLHCYPLESFVDQWPSNPPAFRQVVGTYSTEARALALRLLEAISESL : 180

\*                      200                      \*                      220                      \*                      240  
 LpF3Ha : GLERGHMVKAMGRHAQHMAVNYYPPCPQPELTYGLPGHTDPNALTILLMDPHVSGLQVLR : 240

\*                      260                      \*                      280                      \*                      300  
 LpF3Ha : DGAKWIAVHPRPNALVINLGDQLQALSNGAYKSVWHRAVVNAEQERLSVASFLCPCNSAV : 300

\*                      320                      \*                      340  
 LpF3Ha : ICPAPRLVGDEDPVYRSYTYDEYYKRFWSRNLDQEHCLELFRSQH : 346

## FIGURE 108

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```

      *           20           *           40           *           60
LpF3Ha1 : TCTCNAGACACACTGTGTAACCCAGGTAGCGAGTGGCAAGACTAGCAGAAAGTACGGACA : 60
LpF3Ha2 : --TCNAGACACACTGTGTAACCCAGGTAGCGAGTGGCAAGACTAGCAGAAAGTACGGACA : 58
LpF3Ha3 : ----- : -

      *           80           *           100          *           120
LpF3Ha1 : TCAGCTAACCATTTCCTCAACTAGAATAAGCATGGCTCCGGCGATGTCCAACCCTCTCCTC : 120
LpF3Ha2 : TCAGCTAACCATTTCCTCAACTAGAATAAGCATGGCTCCGGCGATGTCCAACCCTCTCCTC : 118
LpF3Ha3 : ----- : -

      *           140          *           160          *           180
LpF3Ha1 : AGTGATCGGGTGGCAGCGCTCCAAGAAAGTCCCATCTAGCCACGTTAGAGCGGTGGGAGAC : 180
LpF3Ha2 : AGTGATCGGGTGGCAGCGCTCCAAGAAAGTCCCATCTAGCCACGTTAGAGCGGTGGGAGAC : 178
LpF3Ha3 : ----- : -

      *           200          *           220          *           240
LpF3Ha1 : CGCCCAGACCTCGCCAATGTCGACCACGAGTCCGGCGCGGGCATTCCGCTCATCGACCTG : 240
LpF3Ha2 : CGCCCAGACCTCGCCAATGTCGACCACGAGTCCGGCGCGGGCATTCCGCTCATCGACCTG : 238
LpF3Ha3 : ----- : -

      *           260          *           280          *           300
LpF3Ha1 : AAGCAGCTCGAAGGTCCAGGGCGCCGCGAGGGTCGTTCGAGGCCATCGGCTCCGCGTGCGAG : 300
LpF3Ha2 : AAGCAGCTCGAAGGTCCAGGGCGCCGCGAGGGTCGTTCGAGGCCATCGGCTCCGCGTGCGAG : 298
LpF3Ha3 : ----- : -

      *           320          *           340          *           360
LpF3Ha1 : AACGATGGGTTTTTCATGGTGACGAATCATGGCATCCCAGAGGCGGTTCGTGGAGGGGATG : 360
LpF3Ha2 : AACGATGGGTTTTTCATGGTGACGAATCATGGCATCCCAGAGGCGGTTCGTGGAGGGGATG : 358
LpF3Ha3 : ----- : -

      *           380          *           400          *           420
LpF3Ha1 : CTGAGCGTGGCGAGGGAGTTCTTCCACCTGCCGGAGTCCGAGCGGCTCAAGTGCTACTCC : 420
LpF3Ha2 : CTGAGCGTGGCGAGGGAGTTCTTCCACCTGCCGGAGTCCGAGCGGCTCAAGTGCTACTCC : 418
LpF3Ha3 : ----- : -

      *           440          *           460          *           480
LpF3Ha1 : GACGACCCCAAGAAGGCGGTCCGGCTGTTCGACGAGCTTCAACGTGCGCACGGAGAAGGTG : 480
LpF3Ha2 : GACGACCCCAAGAAGGCGGTCCGGCTGTTCGACGAGCTTCAACGTGCGCACGGAGAAGGTG : 478
LpF3Ha3 : ----- : -

      *           500          *           520          *           540
LpF3Ha1 : AGCAACTGGCGCGACTTCCTCCGGCTGCATTGCTACCCTCTTGAGAGCTTCGTCGACCAG : 540
LpF3Ha2 : AGCAACTGGCGCGACTTCCTCCGGCTGCATTGCTACCCTCTTGAGAGCTTCGTCGACCAG : 538
LpF3Ha3 : ----- : -

      *           560          *           580          *           600
LpF3Ha1 : TGGCCGTGGAACCCGCCCGCCTTCAGGCAAGTCGTTCGGCACCTACTCGACGGAAGCGAGA : 600
LpF3Ha2 : TGGCCGTGGAACCCGCCCGCCTTCAGGCAAGTCGTTCGGCACCTACTCGACGGAAGCGAGA : 598
LpF3Ha3 : -----CGGGAAGTCCGC : 12

      *           620          *           640          *           660
LpF3Ha1 : GCGCTGGCGCTGAGGCTCCTTGGAGGCGATATCGGAGAGCCTAGGGCTGGAGAGAGGCCAC : 660
LpF3Ha2 : GCGCTGGCGCTGAGGCTCCTTGGAGGCGATATCGGAGAGCCTAGGGCTGGAGAGAGGCCAC : 658
LpF3Ha3 : CTGCTGGCGCTTCGGCTTCTGGCGCGCATCTCGCTGGGCTGGGGCTGGACNGAGCGTAT : 72

      *           680          *           700          *           720
LpF3Ha1 : ATGGTGAAGGCCATGGGGCGGCACGCGCAGCACATGGCGGTGAACTACTACCCGCCGTGC : 720
LpF3Ha2 : ATGGTGAAGGCCATGGGGCGGCACGCGCAGCACATGGCGGTGAACTACTACCCGCCGTGC : 718
LpF3Ha3 : CTGCGAAGGTCCTGGGCGAGCAGCGAGCAGCACATGGCGGTGAACTACTACCCGCCGTGC : 132

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FIGURE 109

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      *           740           *           760           *           780
LpF3Ha1 : CCGCAGCCGGAGCTCACCTACGGTCTGCCAGGGCA----- :755
LpF3Ha2 : CCGCAGCCGGAGCTCACCTACGGTCTGCCAGGGCACACGGACCCCAATGCCCTCACCATN :778
LpF3Ha3 : CCGCAGCCGGAGCTCACCTACGGTCTGCCCAAGCACACGGACCCCAACGCCCTCACCATC :192

      *           800           *           820           *           840
LpF3Ha1 : ----- : -
LpF3Ha2 : CT----- :780
LpF3Ha3 : CTCCTCATGGATCCCCACGTCTCCGGCCTCCAGGTCTCAGGGACGGCGCCAAGTGGATC :252

      *           860           *           880           *           900
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : GCCGTCCACCCACGCCCCAACGCCCTGGTCATCAACCTAGGCGACCAGCTACAGGCGCTG :312

      *           920           *           940           *           960
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : AGCAACGGCGCGGTACAAGAGCGGTGTGGCACCAGGGCAGTGGTGAACGCGGAGCAGGAGCGT :372

      *           980           *           1000           *           1020
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : CTGTCCGGTGGCATCTTTCCCTGTGCCCGTGCAACAGCGCGGTTATCTGCCCGCGCCGAGG :432

      *           1040           *           1060           *           1080
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : CTCGTCCGCCGACGGGGAGGACCCCGTCTACCGGAGCTACACCTACGACGAGTACTACAAG :492

      *           1100           *           1120           *           1140
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : AGGTTTTTGGAGCAGGAACCTGGATCAGGAGCACTGCCTCGAGCTCTTCAGGAGTCAGCAC :552

      *           1160           *           1180           *           1200
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : TGATGCTTGAACCTTGAGTTACTAGCTAGCTCTCCTTAACAGTGCAAATCCATGGCCCAA :612

      *           1220           *           1240           *           1260
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : GAGGGCCCCGATTGCATGGTTACTTATGTTGTTTGAAGTGGTATTGCTTAACTGCCTAAT :672

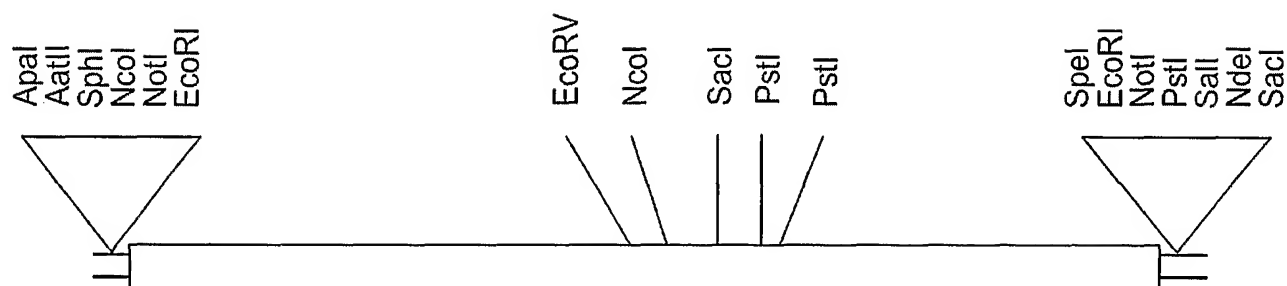
      *           1280           *           1300           *           1320
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : AACATTGCTACATTCTACTNCTATCTTGTCGCTTTAAAATTATAAGATGGCCTAACCTTT :732

      *           1340           *           1360           *           1380
LpF3Ha1 : ----- : -
LpF3Ha2 : ----- : -
LpF3Ha3 : TTCTTAATTGTATGCATNCTGAACATATTTAAGTGTGTGTGTTTCAGACAGTTTAGTCTGC :792

LpF3Ha1 : - : -
LpF3Ha2 : - : -
LpF3Ha3 : A : 793

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FIGURE 109 (cont)

**183/271****LpF3OH****FIGURE 110**

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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GACTCTCAGA
51 ACACACTGTG TAACCACGGT AGCGAGTGGC AAGACTAGCA GAAAGTACGG
101 ACATCAGCTA ACCATTCCTC AACTAGAATA AGCATGGCTC CGGCGATGTC
151 CAACCCTCTC CTCAGTGATC GGGTGGCACG CTCCAAGAAA GTCCCATCTA
201 GCCACGTTAG AGCGGTGGGA GACCGCCAG ACCTCGCCAA TGTCGACCAC
251 GAGTCCGGCG CGGGCATTCC GCTCATCGAC CTGAAGCAGC TCGAAGGTCC
301 AGGGCGCCGC AGGGTCGTCG AGGCCATCGG CTCCGCGTGC GAGAACGATG
351 GGTTTTTTCAT GGTGACGAAT CATGGCATCC CAGAGGCGGT CGTGGAGGGG
401 ATGCTGAGCG TGGCGAGGGA GTTCTTCCAC CTGCCGGAGT CGGAGCGGCT
451 CAAGTGCTAC TCCGACGACC CCAAGAAGGC GGTCCGGCTG TCGACGAGCT
501 TCAACGTGCG CACGGAGAAG GTGAGCAACT GGCGCGACTT CCTCCGGCTG
551 CATTGCTACC CTCTTGAGAG CTTCGTCGAC CAGTGGCCGT CGAACCCGCC
601 CGCCTTCAGG CAAGTCGTCG GCACCTACTC GACGGAAGCG AGAGCGCTGG
651 CGCTGAGGCT CCTGGAGGCG ATATCGGAGA GCCTAGGGCT GGAGAGAGGC
701 CACATGGTGA AGGCCATGGG GCGGCACGCG CAGCACATGG CGGTGAACTA
751 CTACCCGCCG TGCCCGCAGC CGGAGCTCAC CTACGGTCTG CCAGGGCACA
801 AGGACCCCAA TGCCATCACG CTCCTCCTGC AGGACGGCGT CTCCGGCCTG
851 CAGGTCCAGC GCGACGGCCG GTGGGTGGCC GTCAACCCGG TGCCCAACGC
901 CCTCGTCATC AACATCGGCG ATCAGTTACA GGCCTGAGC AACGACCGAT
951 ACAAGAGCGT GAACCACAGA GTGATCGTCA ACAGCGCGAG CGAGAGGATT
1001 TCGGTGCCGA CGTTCTACTG CCCGTCGCCG GACACGGTGG TCGCGCCGGC
1051 CGACGCGCTG GTGGACGACG CCCACCCCTCG GGCTTACCAG CCCTTCACGT
1101 ACCAGGAGTA CTACGAGGAG TTCTGGAAGA TGGGCCTTCA GTCAGCAAGT
1151 TGCTTCGACA GGTTCCGACG GATCGAGTGA TGGACAAGAC GTGGGCCGTT
1201 GTTATCTCCT GGGCCATGAG CGTTGCCGCA GCCGATGTGT CGCCATATGG
1251 TGGAGACGTT TCCTCCCTCC GGAAAAGAAA AATAAAACAG AGTGGAGACC
1301 ACTAGAACCG TCAGATAGCA TCCCAAAAAA AAAAAAAAAA AAAAAAAAAA
1351 AAAAGTACTC TGCGTTGTTA CCACTGCTTA ATCACTAGTG AATTC

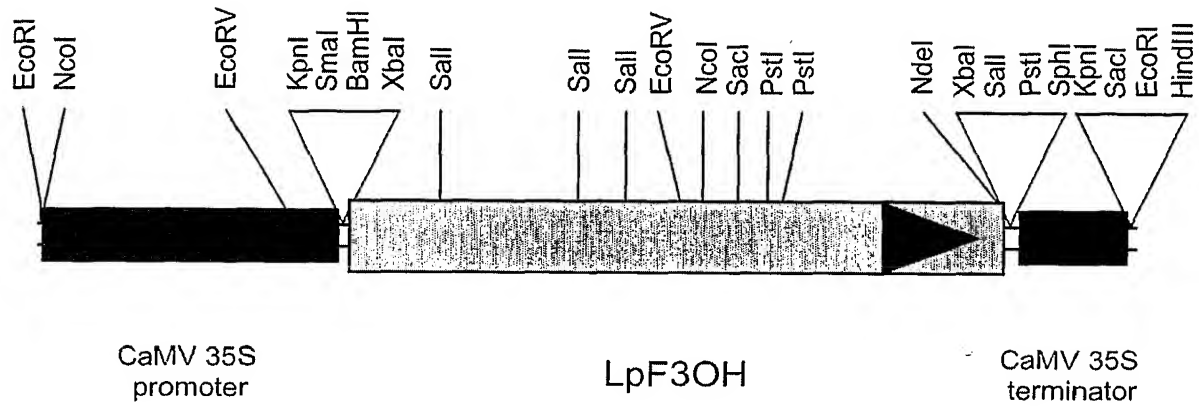
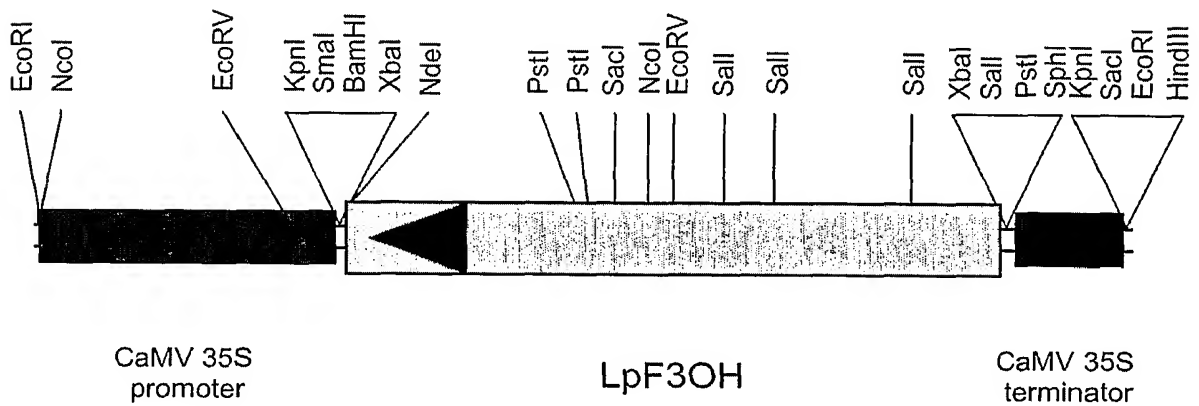
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## FIGURE 111

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1 MAPAMSNPLL SDRVARSKKV PSSHVRAVGD RPDLANVDHE SGAGIPLIDL  
51 KQLEGPGRRR VVEAIGSACE NDGFFMVTNH GIPEAVVEGM LSVAREFFHL  
101 PESERLKCYS DDPKKAVRLS TSFNV RTEKV SNWRDFLR LH CYPLESFVDQ  
151 WPSNPPAFRQ VVGTYSTEAR ALALRLLEAI SESLGLERGH MVKAMGRHAQ  
201 HMAVNYYPFC PQPELTYGLP GHKDPNAITL LLQDGVSG LQ VQRDGRWVAV  
251 NPVPNALVIN IGDQLQALSN DRYKSVNHRV IVNSASERIS VPTFYCPSPD  
301 TVVAPADALV DDAHPRAYQP FTYQEYYEEF WKMGLQSASC LDRFRRIE

## FIGURE 112

**186/271****pDH51LpF3OH sense****pDH51LpF3OH anti****FIGURE 113**



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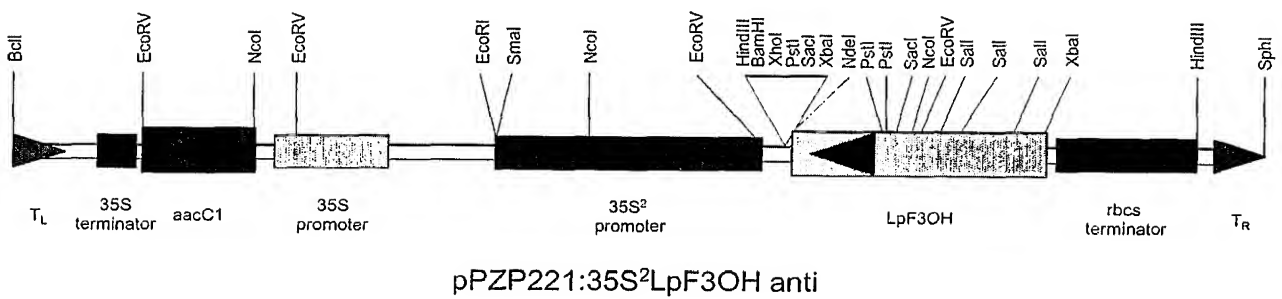
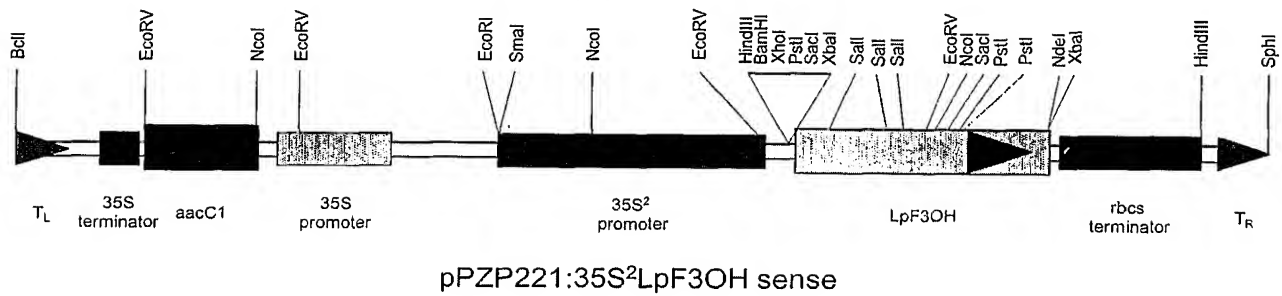


FIGURE 114

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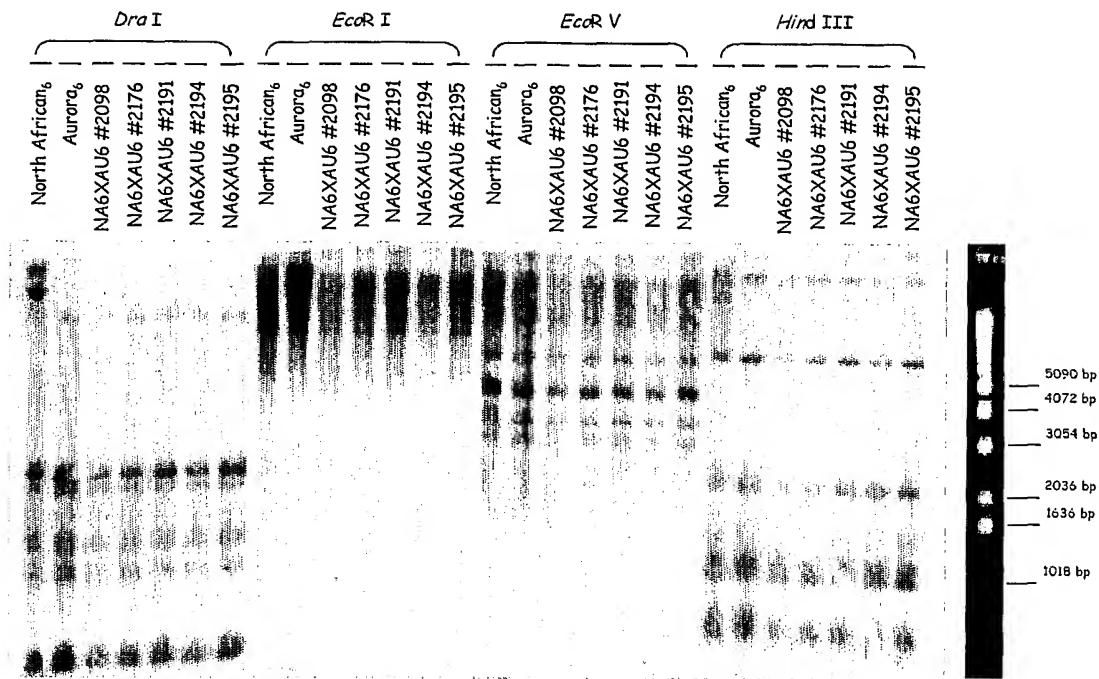
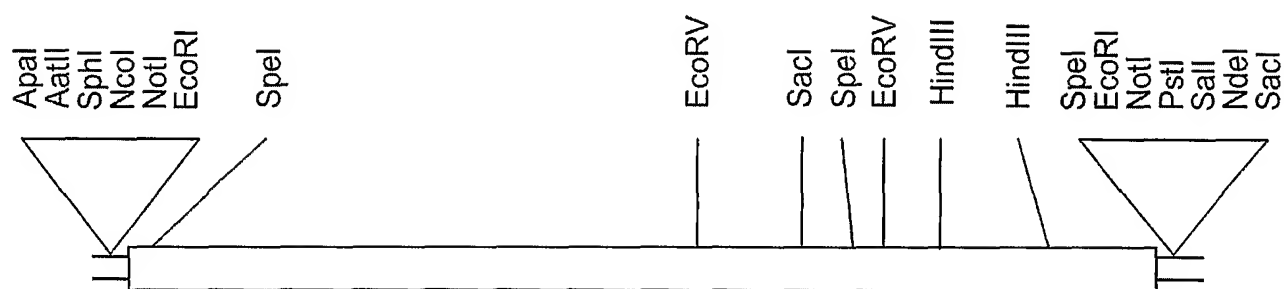


FIGURE 115

**189/271****TrBANa****FIGURE 116**

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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG ATAAAAACTG
51  CACTAGTGTG TATAAGTTTC TTGGTGAAAA AAGAGTTTGT AAATTAACAT
101 CATGGCTAGT ATCAAACAAA TTGGAAACAA GAAAGCATGT GTGATTGGTG
151 GCACTGGTTT TGTTGCATCT ATGTTGATCA AGCAGTTACT TGAAAAGGGT
201 TATGCTGTTA ATACTACCGT TAGAGACCCA GATAGCCCTA AGAAAATATC
251 TCACCTAGTG GCACTGCAAA GTTTGGGGGA ACTGAATCTA TTTAGAGCAG
301 ACTTAACAGT TGAAGAAGAT TTTGATGCTC CTATAGCAGG ATGTGAACTT
351 GTTTTTCAAC TTGCTACACC TGTGAACTTT GCTTCTCAAG ATCCTGAGAA
401 TGACATGATA AAGCCAGCAA TCAAAGGTGT GTTGAATGTG TTGAAAGCAA
451 TTGCAAGAGC AAAAGAAGTT AAAAGAGTTA TCTTAACATC TTCGGCAGCC
501 GCGGTGACTA TAAATGAACT CAAAGGGACA GGTCATGTTA TGGATGAAAC
551 CAACTGGTCT GATGTTGAAT TTCTCAACAC TGCAAAACCA CCCACTTGGG
601 GTTATCCTGC CTCAAAAATG CTAGCTGAAA AGGCTGCATG GAAATTTGCT
651 GAAGAAAATG ACATTGATCT AATCACTGTG ATACCTAGTT TAACAACCTGG
701 TCCTTCTCTC ACACCAGATA TCCCATCTAG TGTGCGCTTG GCAATGTCTC
751 TAATAACAGG CAATGATTTT CTCATAAATG CTTTGAAAGG AATGCAGTTT
801 CTGTCGGGTT CGTTATCCAT CACTCATGTT GAGGATATTT GCCGAGCTCA
851 TATATTTCTT GCAGAGAAAG AATCAGCTTC TGGTAGATAC ATTTGCTGTG
901 CTCACAATAC TAGTGTTCCC GAGCTTGCAA AGTTTCTCAA CAAACGATAT
951 CCTCAGTATA AAGTTCCAAC TGAATTTGAT GATTGCCCCA GCAAGGCAAA
1001 GTTGATAATC TCTTCTGAAA AGCTTATCAA AGAAGGGTTC AGTTTCAAGC
1051 ATGGTATTGC CGAAACTTTC GACCAGACTG TCGAGTATTT TAAGACTAAG
1101 GGGGCACTGA AGAATTAGAT TTTGATATTT CTAATTCAAT AGCAAACCTCT
1151 AAGCTTGTTA TGTGTTTGTG AAGTTCAGAG TGAAATATCA AATGAATAAG
1201 TGGAGAGAGC ACAATAAGAG GAGAGCACAA TAATTTTGGA AAAAAAAAAA
1251 AAAAAAAAAA AAAAAAAGT ACTCTGCGTT GTTACCACTG CTTAATCACT
1301 AGTGAATTC

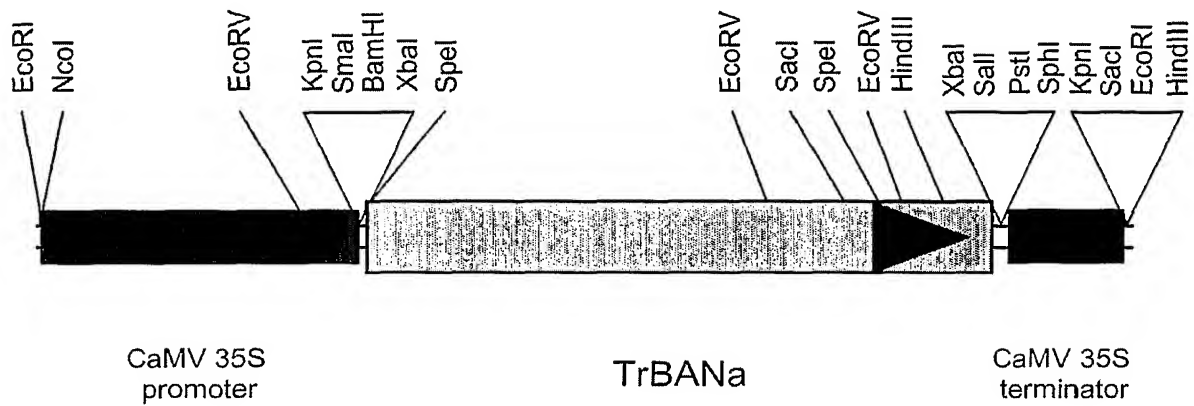
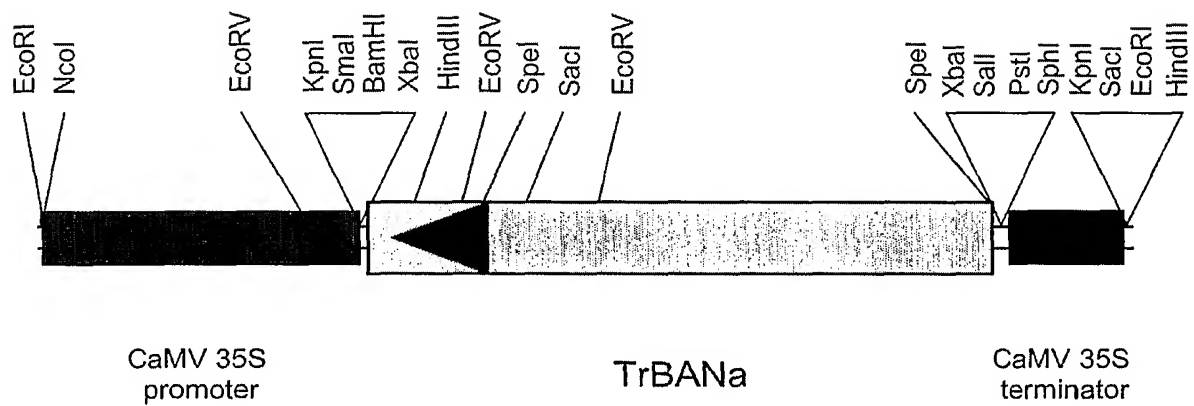
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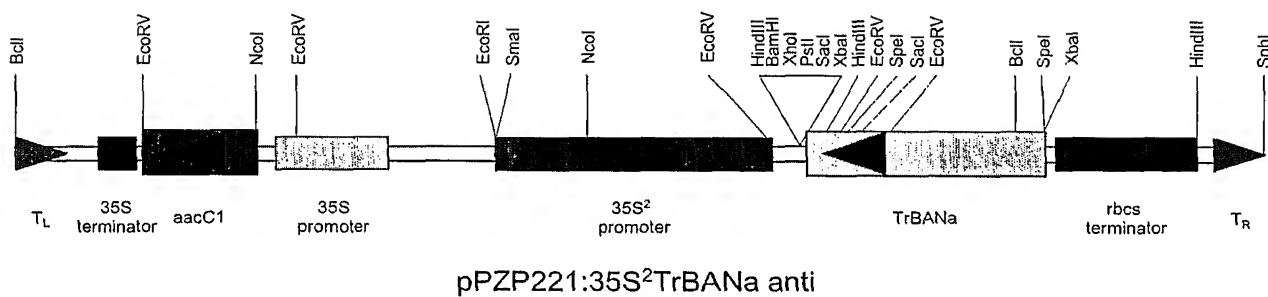
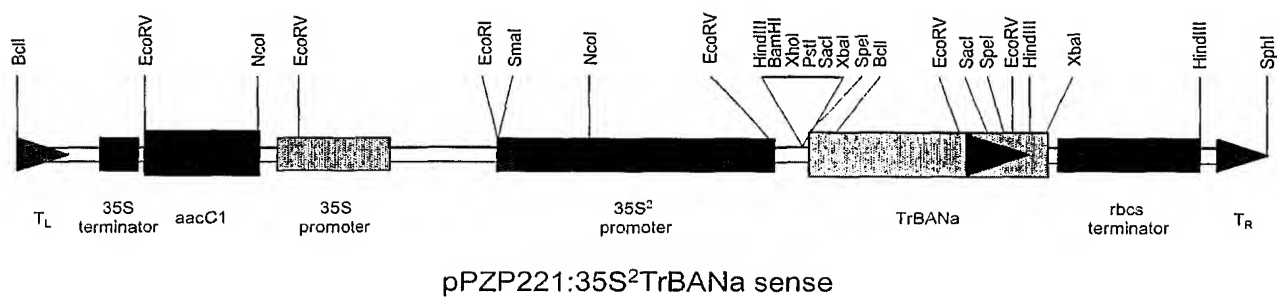
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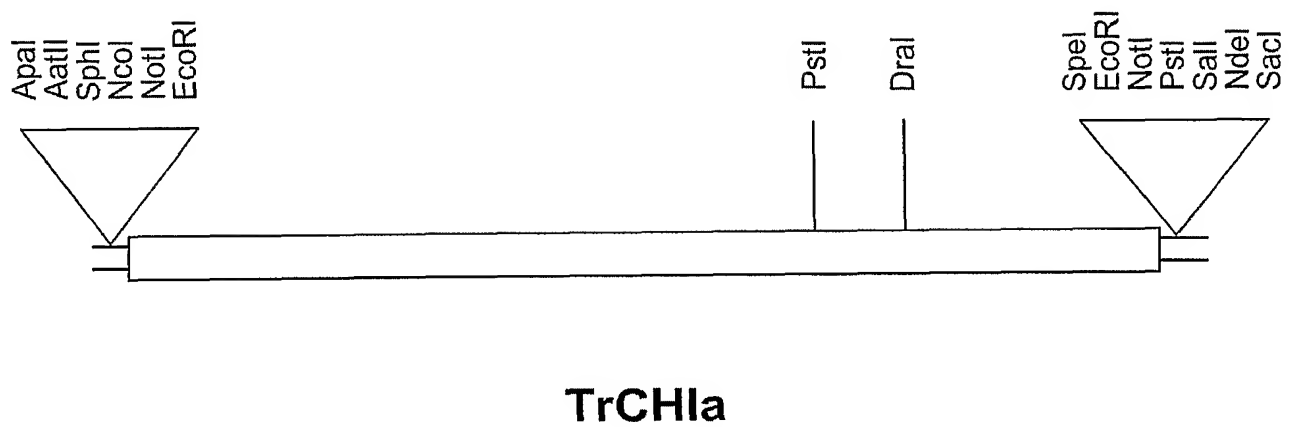
# 191/271

1 MASIKQIGNK KACVIGGTGF VASMLIKQLL EKGAVNTTV RDPDSPKKIS  
51 HLVALQSLGE LNLFRADLTV EEDFDAPIAG CELVFQLATP VNFASQDPEN  
101 DMIKPAIKGV LNVLKAIARA KEVKRVILTS SAAAVTINEL KGTGHVMDDET  
151 NWSDV EFLNT AKPPTWGYP A SKMLAEKAAW KFAEENDIDL ITVIPSLLTTG  
201 PSLTPDIPSS VGLAMSLITG NDFLINALKG MQFLSGSLSI THVEDICRAH  
251 IFLAEKESAS GRYICCAHNT SVPELAKFLN KRYPQYKVPT EFDDCPSKAK  
301 LIISSEKLIK EGFSFKHGIA ETFDQTV EYF KTKGALKN

## FIGURE 118

**192/271****pDH51TrBANa sense****pDH51TrBANa anti****FIGURE 119**

**193/271****FIGURE 120**

**194/271****FIGURE 121**



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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GACTTAAACA
51 TTGACACAAG TCCCAAATAA AAAAGATCTG AAACAACATA GTCACCCCAT
101 TTTTAAACAT TAAACTAAAA ATATGTCGGC CATCACCGCA ATCCAAGTCG
151 AGAACCTTGA ATTTCCGGCT GTGGTTACTT CTCCGGCCAC CGGTAAGTCA
201 TATTTTCTTG GTGGTGCAGG GGAGAGAGGT TTGACTATTG AAGGAACTT
251 CATCAAGTTC ACTGCCATAG GAGTATATTT GGAAGATGTA GCAGTGGCTT
301 CACTTGCCAC TAAATGGAAG GGTAAATCCT CTGAGGAGTT GCTTGAGACT
351 CTTGACTTCT ATAGAGACAT CATTTTCAGGA CCCTTTGAAA AGTTGATTCTG
401 AGGATCGAAG ATTAGGGAAT TGAGTGGTCC TGAGTACTCA AGGAAGGTTA
451 ATGAAAAC TGCGGCACAC TTAAAATCTG TTGGGACTTA TGGAGATGCT
501 GAAGCTGAAG CTATGCAAAA ATTTGTTGAA GCCTTCAAGC CTATTAATTT
551 TCCACCTGGT GCCTCTGTTT TTTACAGGCA ATCACCTGAT GGAATATTAG
601 GGCTTAGTTT CTCTCAAGAT GCAAGTATAC CAGAAAAGGA GGCTGCAGTA
651 ATAGAGAACA AGGCAGCTTC ATCGGCAGTG TTAGAAACTA TGATTGGTGA
701 ACATGCTGTT TCTCCTGATT TAAAGCGTTG TTTGGCTGCA AGATTACCTG
751 CTTGTTGAA CGAGGGTACT TTCAAGATTG AATGAAAAC TATTATTATT
801 ATCTCCAAAA GCATTGCAGC ACAAGATTGA GTCATTTATG AGCATGGACA
851 TTTTATATGTC CACACATGTT TAACTTTTGT ATCTCTCTTT AGATTCTCAT
901 CAATATCAAT AATACTAATA TGAAACGAAG TCAAAAAAAA AAAAAAAAAA
951 AAAAAAAAAA AAAAGTACTC TCGTTGTTA CCACTGCTTA ATCACTAGTG
1001 AATTC

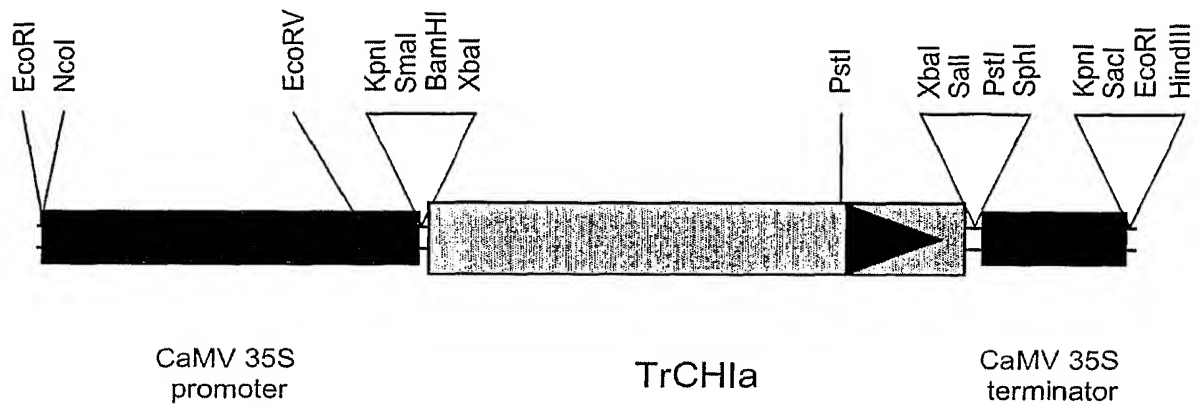
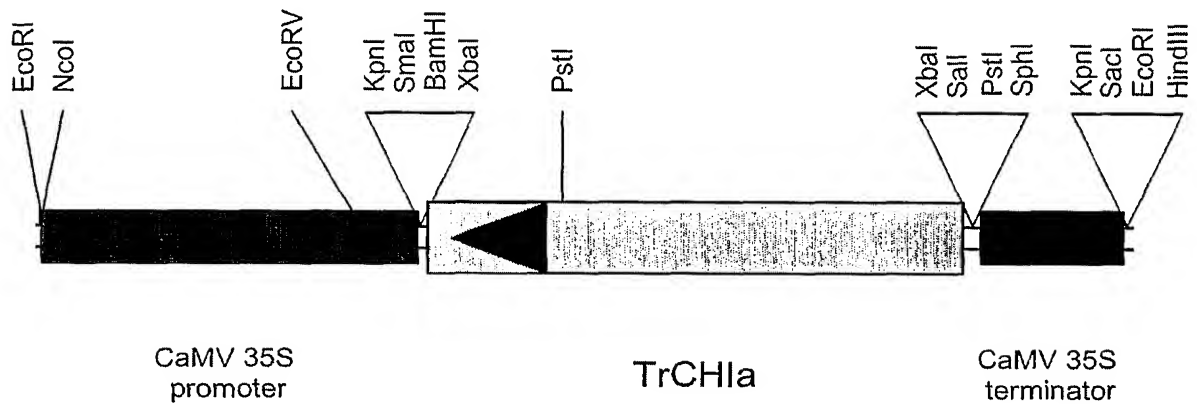
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## FIGURE 122

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1 MSAITAIQVE NLEFPAVVTs PATGKSYFLG GAGERGLTIE GNFIKFTAIG  
51 VYLEDVAVAS LATKWKGKSS EELLETLDFY RDIISGPFEK LIRGSKIREL  
101 SGPEYSRKVN ENCV AHLKSV GTYGDAEAEA MQKFVEAFKP INFPPGASVF  
151 YRQSPDGILG LSFSQDASIP EKEAAVIENK AASSAVLETM IGEHAVSPDL  
201 KRCLAARLPA LLNEGTFKIE

## FIGURE 123

**197/271****pDH51TrCHla sense****pDH51TrCHla anti****FIGURE 124**

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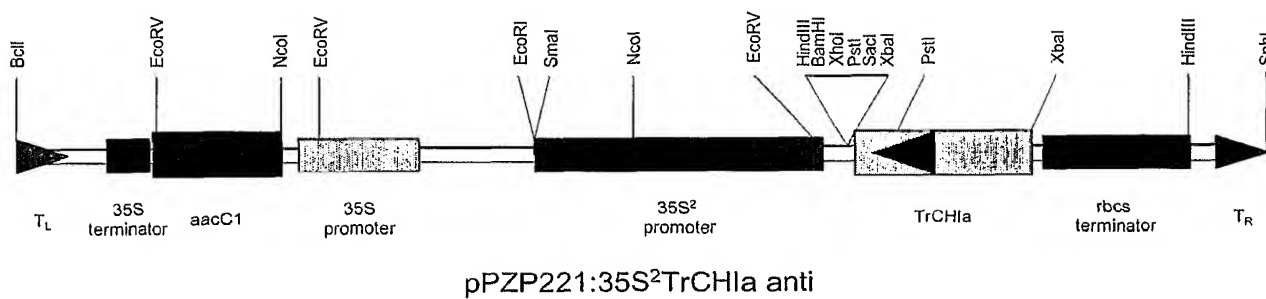
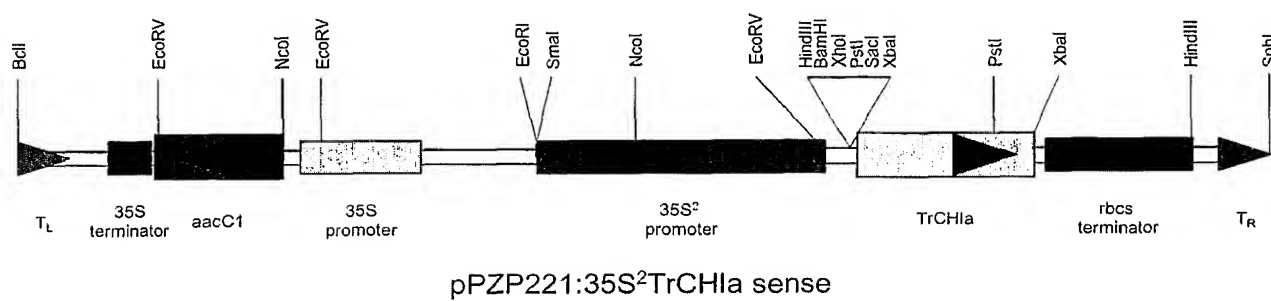
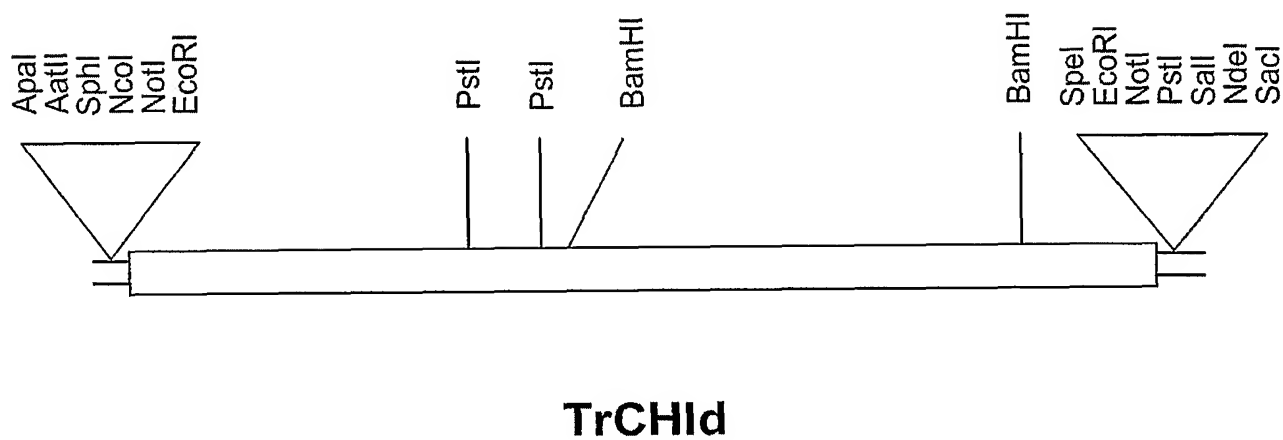


FIGURE 125

**199/271****FIGURE 126**

# 200/271

```

1  GAATTCACTA GTGATTAAGC AGTGGTAACA ACGCAGAGTA CGCGGGGACA
51 TTACAACTCA CAACACCTTC TCCATTACCA TCTATCTTCT ACTAAGTTCA
101 ACGAGATCAA TGGCACTTCC TTCTGTCACC GCTTTGAATA TCGAGAACAA
151 TCTATTCCCT CCTACCGTCA CACCACCGGG ATCCACCAAC AATTTCTTCC
201 TCGGCGGTGC AGGAGAGCGG GGTCTTCAAA TTCAAGACAA ATTTGTCAAA
251 TTCACCGCTA TTGGTGTTTA TCTACAGGAC ATTGCTGTTC CTTACCTCGC
301 CACTAAATGG AAGGGTAAGA CTGCTCAAGA GCTAACGGAA ACTGTTCTTT
351 TCTTCAGGGA CATCGTTACA GGTCCATTTC AGAAATTTAT GCAGGTGACA
401 ATGATCTTGC CATTGACTGG GCAACAATAC TCAGAGAAAG TGTCAAGAAA
451 TTGTGTAGCT ATTTGGAAGT CTCTTGGGAT TTATACCGAC GAAGAAGCCA
501 AAGCAATTGA GAAGTTTGTT TCTGTCTTCA AAGATGAAAC ATTCCCACCA
551 GGCTCCTCTA TCCTTTTCAC AGTATTACCC AAAGGATTAG GATCACTAAC
601 GATAAGTTTC TCTAAAGATG GATCCATTCC AGAGACCGAG TCTGCAGTTA
651 TAGAGAATAA GCTACTCTCA CAAGCTGTGC TTGAGTCGAT GATAGGGGCG
701 CACGGTGTCT CCCCTGCAGC AAAACAGAGT TTGGCCACCA GGTATCCGA
751 GTTATTCAAC GAGGTTGGTG ATGCTAGCAA CTGATTATAT CAACAAAACG
801 AAAATGAAAG TCCTTTCTGC AATAAAGACC AAGCGGAAAT TTTATTTTAG
851 GTGCACTTTG AAATGACCTC TTTGGCGACT TTTTCTTGTA CTAATAATAA
901 AGAGTGTGTT TGTATCATGT TGTAATTTTA TTTTAGAAAA AGTGAGGTAA
951 GAAAGGAGTC CTTATGTTTA TTTCAATTAT TGAAAAATTA TTTGCATGTA
1001 TAATTGATTT CAACTGATGT TATTTAATCA CGTTTTTTCT AAAAAAAAAA
1051 AAAAAAAAAA AAAAAAAAAA GTACTCTGCG TTGTTACCAC TGCTTAATCG
1101 AATTC

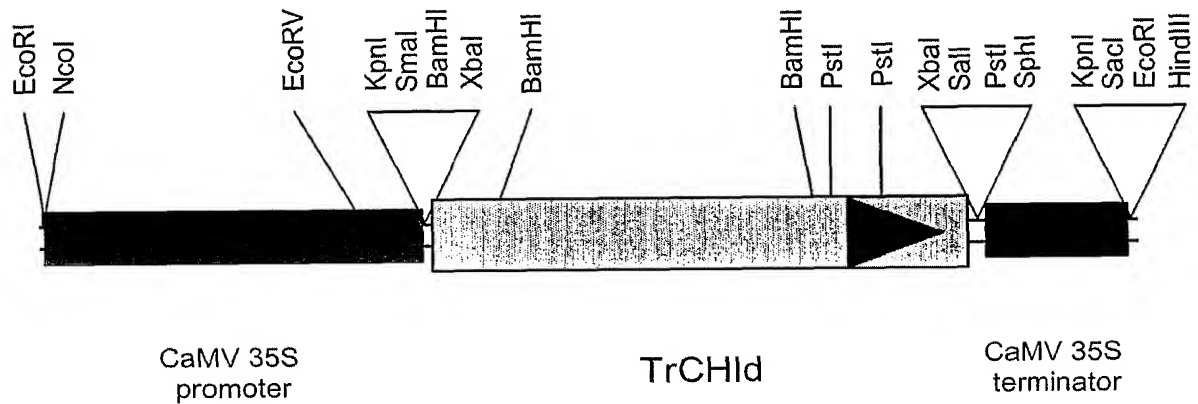
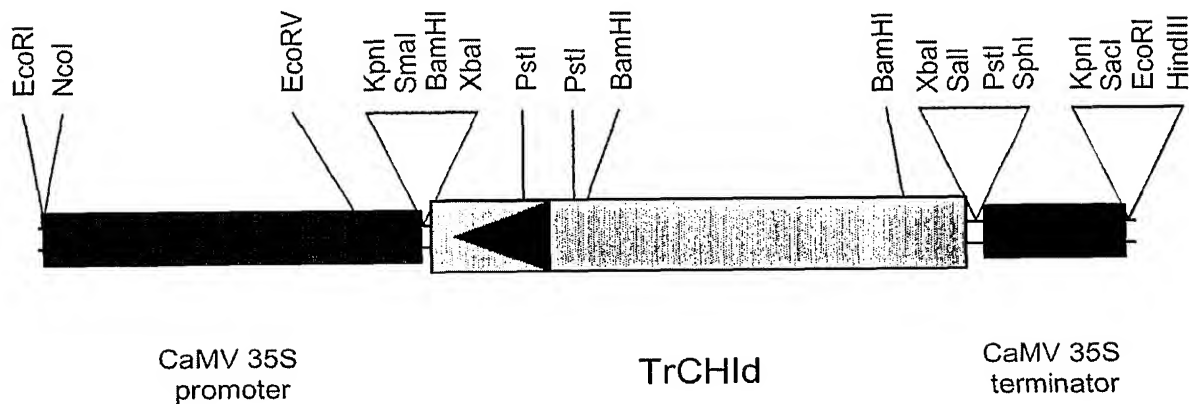
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## FIGURE 127

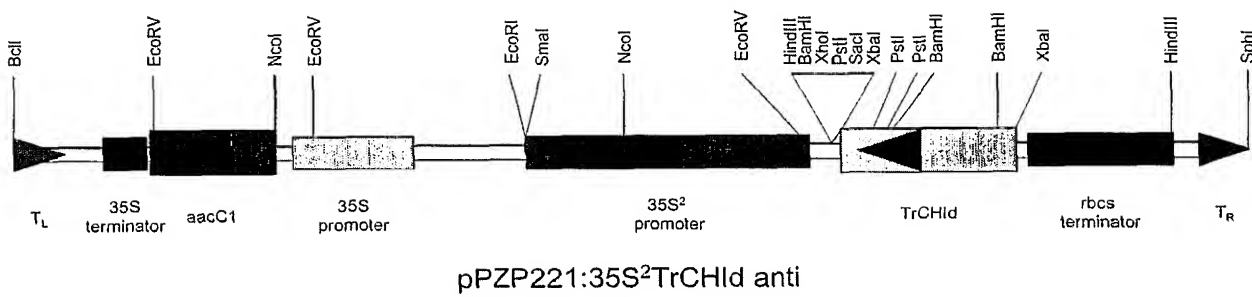
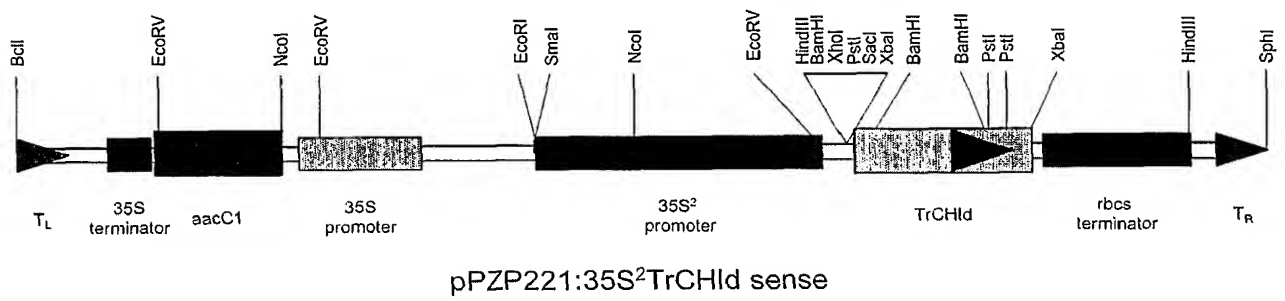
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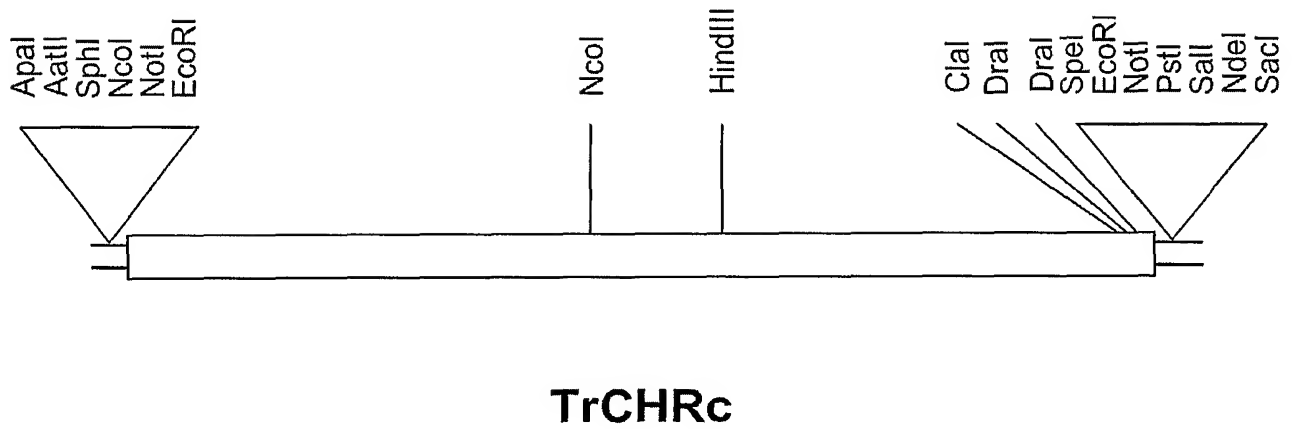
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51 IGVYLQDIAV PYLATKWKGK TAQELTETVP FFRDIVTGPF EKFMQVTMIL  
101 PLTGQQYSEK VSENCVAIWK SLGIYTDEEA KAIEKFVSVF KDETFFPGSS  
151 ILFTVLPKGL GSLTISFSKD GSIPETESAV IENKLLSQAV LESMIGAHGV  
201 SPAAKQSLAT RLSELFNEVG DASN

## FIGURE 128

**202/271****pDH51TrCHId sense****pDH51TrCHId anti****FIGURE 129**



**203/271****FIGURE 130**

**204/271****FIGURE 131**

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```

1  GAATTCGATT AAGCAGTGGT AACCAACGCAG AGTACGCGGG GATTCAAACA
51 TAGCTCAAAG TGTGTAACAA ATTTCTTAAC TTAAAACATT TTCAACCCAA
101 CAAAAAAAAA CAAAGACAAA AACATGGGTA GTGTTGAAAT TCCAACAAAG
151 GTTCTTACTA ACAGTTCTAG TCAAGTGAAA ATGCCTGTGG TTGGAATGGG
201 ATCAGCACCT GATTTCACAT GTAAGAAAGA CACAAAAGAT GCAATCATTG
251 AAGCCATCAA ACAGGGTTAT AGACACTTTG ATACTGCTGC TGCTTATGGC
301 TCAGAACAAG CTCTTGGTGA AGGTTTGAAA GAAGCAATTG AACTTGGTCT
351 TGTCACTAGA GAAGACCTTT TTGTTACTTC TAAACTTTGG GTCACTGAAA
401 ATCATCCTCA TCTTGTTGTT CCTGCTCTTC AAAAATCTCT CAAGACTCTT
451 CAATTGGAGT ACTTGGACTT GTATTTGATC CATTGGCCAC TTAGTTCTCA
501 GCCTGGAAAG TTTTCATTTT CAATTGATGT GGCAGATCTC TTGCCATTTG
551 ATGTGAAGGG TGTTTGGGAA TCCATGGAAG AAGGCTTGAA ACTTGGACTC
601 ACTAAAGCTA TTGGTGTTAG TAACTTCTCT GTCAAGAAAC TTCAAAATCT
651 TGTCTCAGTT GCCACTGTTT TTCCTGCTGT CAATCAAGTG GAGATGAACC
701 TTGCATGGCA ACAAAGAAG CTTAGAGAAT TTTGCAATGC AAATGGAATA
751 GTGTTAACTG CATTTTCACC ATTGAGAAAA GGTGCAAGCA GGGGACCAAA
801 TGAAGTTATG GAAAATGATA TGCTTAAAGA GATTGCAGAT GCTCATGGAA
851 AGTCTGTTGC ACAAATTTCA TTGAGATGGT TATATGAACA AGGAGTCACT
901 TTTGTTCCCA AGAGCTATGA TAAGGAAAGA ATGGGTCAAA ATTTGGCTAT
951 CTTTGATTGG ACATTGGCAA AAGAAGATCA TGAGAAAATT GATCAAATTA
1001 AGCAGAACCG TTTGATCCCT GGACCAACCA AGCCAGGACT CAGTGACCTA
1051 TGGGATGATG AAATATAAAG TGGAAGATGT TAAAAGTCCC TTAAGCTCAC
1101 TCAATATCTA TCTATTGTGT ACTTTTGTCA TTTGGGGTTT GAAATTGAGT
1151 CACCCTTGTT TCTGTATCGA TTTAAAATTT AAATAATCAA TTTTTCATTA
1201 CAAAAAAAAA AAAAAAAAAA AAAAAAAAAA AGTACTCTGC GTTGTTACCA
1251 CTGCTTAATC ACTAGTGAAT TC

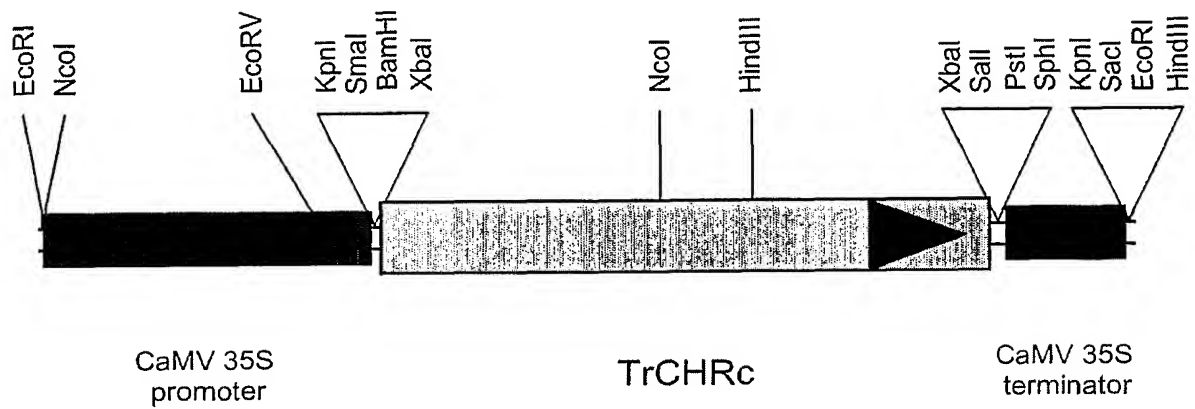
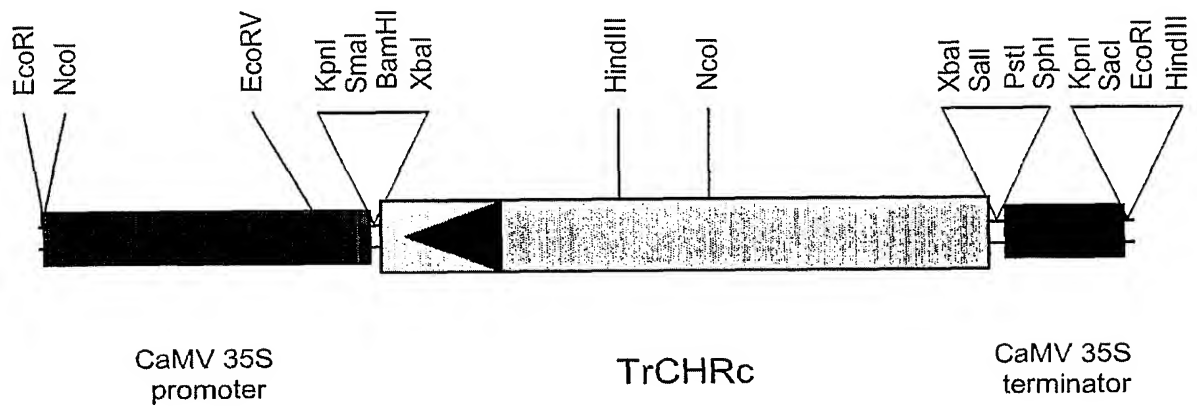
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## FIGURE 132

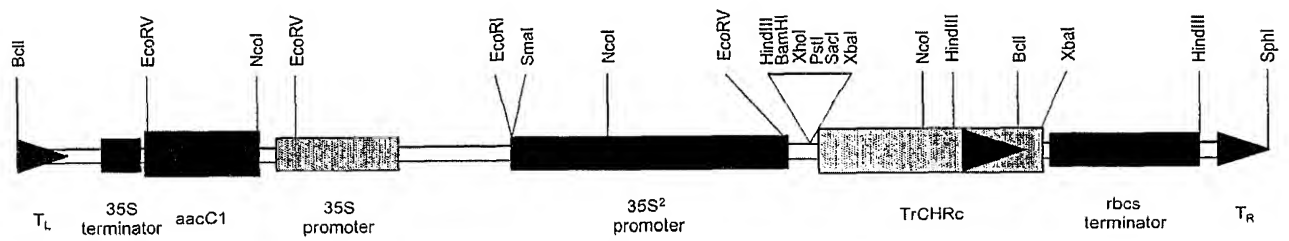
# 206/271

1 MGSVEIPTKV LTNSSSQVKM PVVGMGSAPD FTCKKDTKDA IIEAIKQGYR  
51 HFDTAAAYGS EQALGEGGLKE AIELGLVTRE DLFVTSKLWV TENHPHLVVP  
101 ALQKSLKTLQ LEYLDLYLIH WPLSSQPGKF SFPIDVADLL PFDVKGVWES  
151 MEEGLKLGLT KAIGVSNFSV KKLQNLVSVA TVLPAVNQVE MNLAWQQKKL  
201 REFECNANGIV LTAFSPLRKG ASRGPNEVME NDMLKEIADA HGKSVAQISL  
251 RWLYEQGVTF VPKSYDKERM GQNLAIWDWT LAKEDHEKID QIKQNRLLPG  
301 PTKPGLSDLW DDEI

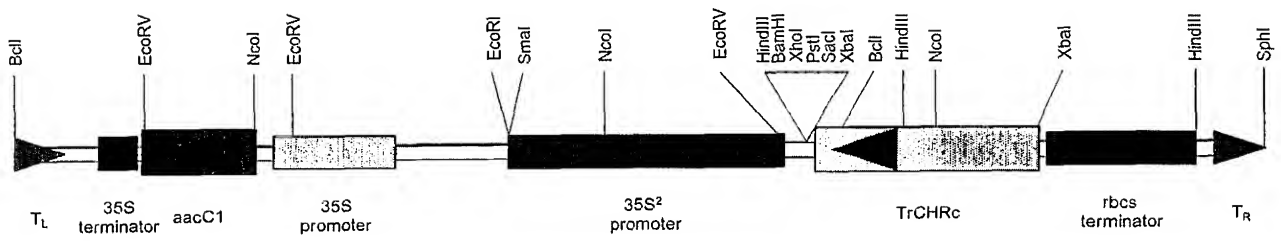
## FIGURE 133

**207/271****pDH51TrCHRC sense****pDH51TrCHRC anti****FIGURE 134**

208/271



pPZP221:35S²TrCHRC sense



pPZP221:35S²TrCHRC anti

FIGURE 135

**209/271****TrCHSa1****FIGURE 136**

# 210/271

```

1  GAATTCGATT AAGCAGTGGT ATCAACGCAG AGTACGCGGG GACAACAACT
51 ATAACTTCCTT GTTATTAACC AATTGAGTTTC AAATTACATA CATAGCAGGA
101 ACTATACTAA AGATATCAAC ATGGTTTAGTG TTTCTGAAAT TCGCAAGGCT
151 CAAAGGGCTG AAGGCCCTGC AACTATTTTTG GCCATTGGTA CTGCAAATCC
201 AGCAAATCGT GTTGACCAGA GTACATATCC TGATTTCTAC TTCAAAATCA
251 CTAACAGTGA GCATAAGGTT GAGCTTAAAG AGAAATTTCA GCGCATGTGT
301 GATAAATCTA TGATCAAGAG CAGATACATG TATCTAACAG AAGAGATTTT
351 GAAAGAAAAT CCTAGTCTTT GTGAATACAT GGCACCTTCA TTGGATGCTA
401 GGCAAGACAT GGTGGTGGTT GAGGTACCTA GACTTGGGAA GGAGGCTGCA
451 GTGAAAGCTA TCAAAGAATG GGGTCAACCA AAGTCAAAGA TTACTCACTT
501 AATCTTTTGC ACCACAAGTG GTGTTGACAT GCCTGGTGCC GATTACCAAC
551 TCACAAAACT CTTAGGTCTT CGCCCATATG TGAAGAGGTA CATGATGTAC
601 CAACAAGGGT GCTTTGCAGG TGGGACGGTT CTTCGTTTGG CCAAGGATTT
651 GGCCGAGAAC AACAAAGGTG CTCGTGTGTT GGTGTGTTGC TCTGAAGTAA
701 CCGCAGTCAC ATTCCGCGGC CCCAGTGACA CTCATTTGGA CAGTCTTGTT
751 GGACAAGCAC TATTCGGAGA TGGAGCTGCT GCACTCATTG TTGGCTCAGA
801 CCCAGTACCA GAAATTGAGA AGCCAATATT TGAGATGGTT TGGACCGCAC
851 AGACAATTGC TCCAGATAGT GAAGGTGCCA TTGATGGTCA TCTTCGTGAA
901 GCTGGACTAA CATTTCATCT TCTTAAAGAT GTTCCTGGGA TTGTCTCAA
951 GAACATTGAT AAGGCATTGG TTGAGGCATT CCAACCATTA AACATCTCTG
1001 ATTACAATTC AATCTTTTGG ATTGCTCATC CAGGTGGTCC TGCAATTCTA
1051 GACCAAGTTG AGATAAAGTT GGGCTTAAAA CCTGAAAAAA TGAAGGCCAC
1101 CAGAGATGTA CTTAGTGAAT ATGGTAACAT GTCAAGTGCA TGTGTATTGT
1151 TCATCTTAGA TGAGATGAGA AAGAAATCGG CTGAAAATGG ACTTAAAAACC
1201 ACAGGAGAAG GACTTGACTG GGGTGTGTTG TTTGGATTTG GGCCCGGACT
1251 TACCATTGAA ACTGTTGTTT TACATAGTGT GGCTATATGA GAATGAGAGA
1301 CTTGATTTGT TTTTATTGTA TTGTATTGTA TTACTTTAAAA TCTTGTTGA
1351 ACCTCCATTT TAAGAATAAA TATGGAGTTC AATATGGACC ATCCTGTTAA
1401 AATAATATAT CGTTAATAGC TATTATTTTA GTGTCTGTTT CTTTTTACTA
1451 AACTATTTTA TTTTAGTATT TGTTTTTGAC CAAAAAAAA AAAAAAAAA
1501 AAAAAAGTA CTCTGCGTTG TTACCACTGC TTAATCACTA GTGAATTC

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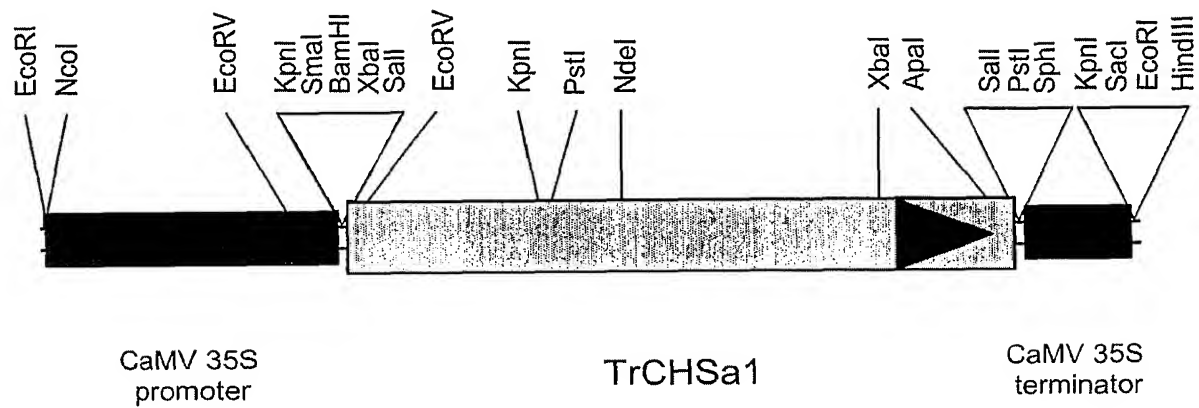
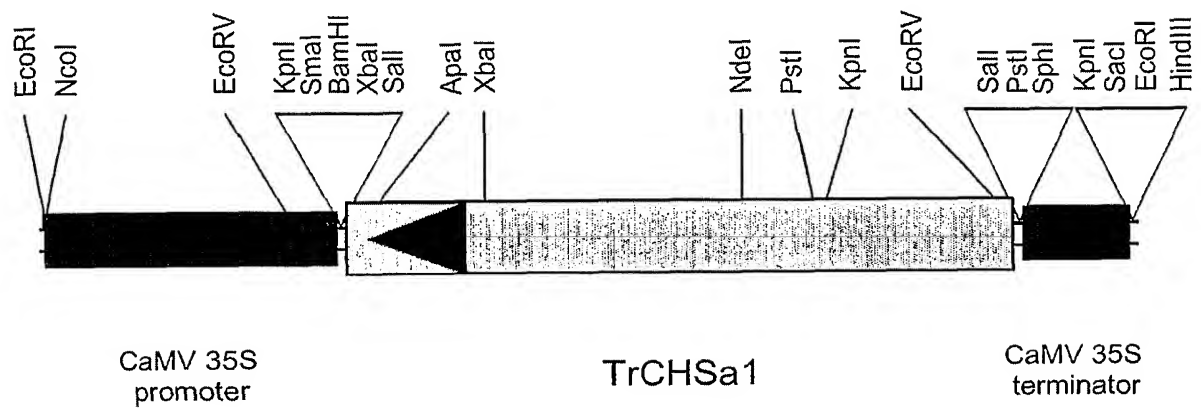
## FIGURE 137

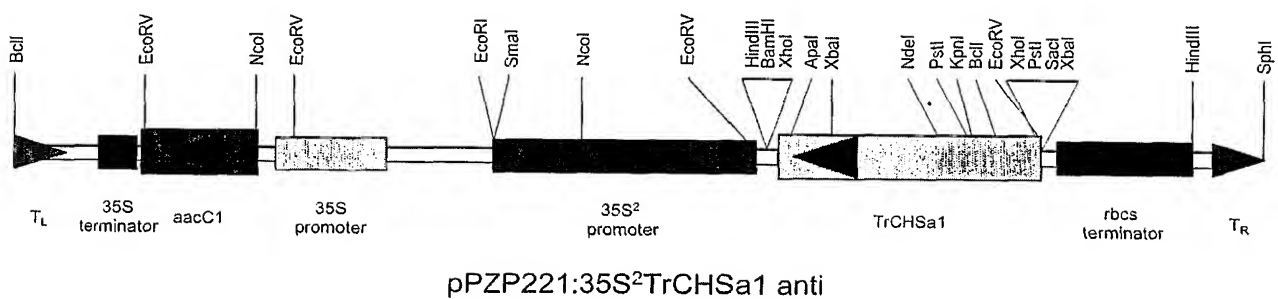
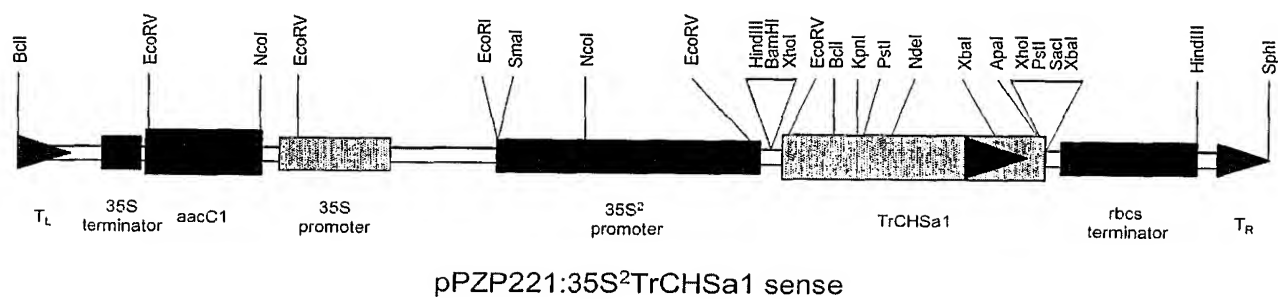


# 211/271

```
1  MVSVSEIRKA QRAEGPATIL AIGTANPANR VDQSTYPDFY FKITNSEHKV
51  ELKEKFQPMC DKSMIKSRYM YLTEEILKEN PSLCEYMAPS LDARQDMVVV
101 EVPRLGKEAA VKAIKEWGQP KSKITHLIFC TTSGVDMPGA DYQLTKLLGL
151 RPYVKRYMMY QQGCFAGGTV LRLAKDLAEN NKGARVLVVC SEVTAVTFRG
201 PSDTHLDSLV GQALFGDGAA ALIVGSDPVP EIEKPIFEMV WTAQTIAPDS
251 EGAIDGHLRE AGLTFHLLKD VPGIVSKNID KALVEAFQPL NISDYNISIFW
301 IAHPPGGPAIL DQVEIKLGLK PEKMKATRDV LSEYGNMSSA CVLFILDEMR
351 KKSAENGLKT TGEGLDWGVL FGFPGPLTIE TVVLHSVAI
```

## FIGURE 138

**212/271****pDH51TrCHSa1 sense****pDH51TrCHSa1 anti****FIGURE 139**

**213/271****FIGURE 140**

**214/271****TrCHSa3****FIGURE 141**

# 215/271

```

1  GAATTCACTA GTGATTAAGC AGTGGTAACA ACGCAGAGTA CGCGGGGAAC
51  AAAAACAAC T ACGCATATTA TATATATATA TATATAGTCT ATAATTGAAA
101 GAAACTGCTA AAGATATTAT TAAGATATGG TGAGTGTAGC TGAAATTCGC
151 AAGGCTCAGA GGGCTGAAGG CCCTGCAACC ATTTTGGCCA TTGGCACTGC
201 AAATCCACCA AACCGTGTTG AGCAGAGCAC ATATCCTGAT TTCTACTTCA
251 AAATTACAAA CAGTGAGCAC AAGACTGAGC TCAAAGAGAA GTTCCAACGC
301 ATGTGTGACA AATCCATGAT CAAGAGCAGA TACATGTATC TAACAGAAGA
351 GATTTTGAAA GAAAATCCTA GTCTTTGTGA ATACATGGCA CCTTCATTGG
401 ATGCTAGGCA AGACATGGTG GTGGTTGAGG TACCTAGACT TGGGAAGGAG
451 GCTGCAGTCA AGGCCATTAA AGAATGGGGT CAACCAAAGT CAAAGATTAC
501 TCACTTAATC TTTTGCACCA CAAGTGGTGT TGACATGCCT GGTGCTGATT
551 ACCAACTCAC AAAACTCTTA GGTCTTCGCC CATATGTGAA AAGGTATATG
601 ATGTACCAAC AAGGTTGTTT TGCAGGAGGC ACGGTGCTTC GTTTGGCAAA
651 AGATTTGGCC GAGAACAACA AAGGTGCTCG TGTGCTAGTT GTTTGTTCTG
701 AAGTCACCGC AGTCACATTT CGCGGCCCCA GTGATACTCA CTTGGACAGT
751 CTTGTTGGAC AAGCATTGTT TGGAGATGGA GCCGCTGCAC TAATTGTTGG
801 TTCTGATCCA GTGCCTGAAA TTGAGAAACC AATATTTGAG ATGGTTTGGA
851 CTGCACAAAC AATTGCTCCA GACAGTGAAG GTGCCATTGA TGGTCATCTT
901 CGTGAAGCTG GGCTAACATT TCATCTTCTT AAAGATGTTC CTGGGATTGT
951 ATCAAAGAAC ATTAATAAAG CATTGGTTGA GGCTTTCCAA CCATTAGGAA
1001 TTTCTGACTA CAACTCAATC TTTTGGATTG CACACCCGGG TGGACCTGCA
1051 ATTCTTGATC AAGTAGAACA AAAGCTAGCC TTGAAGCCCG AAAAGATGAG
1101 GGCCACGAGG GAAGTTCTAA GTGAATATGG AAACATGTCA AGCGCATGTG
1151 TATTGTTTCAT CTTAGATGAG ATGCGGAAGA AATCGGCTCA AAATGGACTT
1201 AAGACAAC TG GAGAAGGACT TGATTGGGGT GTGTTGTTCG GCTTCGGACC
1251 AGGACTTACC ATTGAAACCG TTGTTCTTCG TAGCGTGGCT ATATAAGATG
1301 TGTGATTGTT TTTATTTTAA TGTATTACTT TTAATCTTGC TGCCTTGAAT
1351 TTCGATTTAA GAATAAATAA ATATATCTTT TGATAAAAAA AAAAAAAAAA
1401 AAAAAAAAAA AAGTACTCTG CGTTGTTACC ACTGCTTAAT CGAATTC

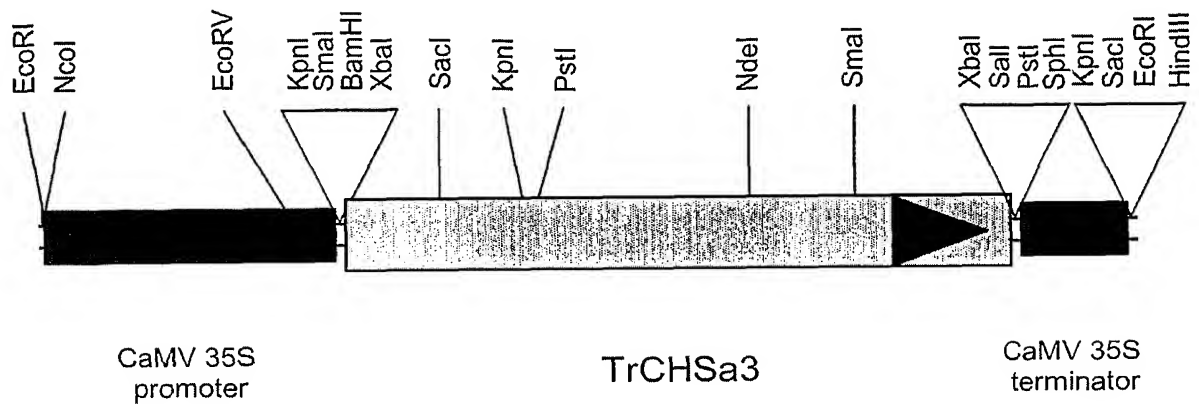
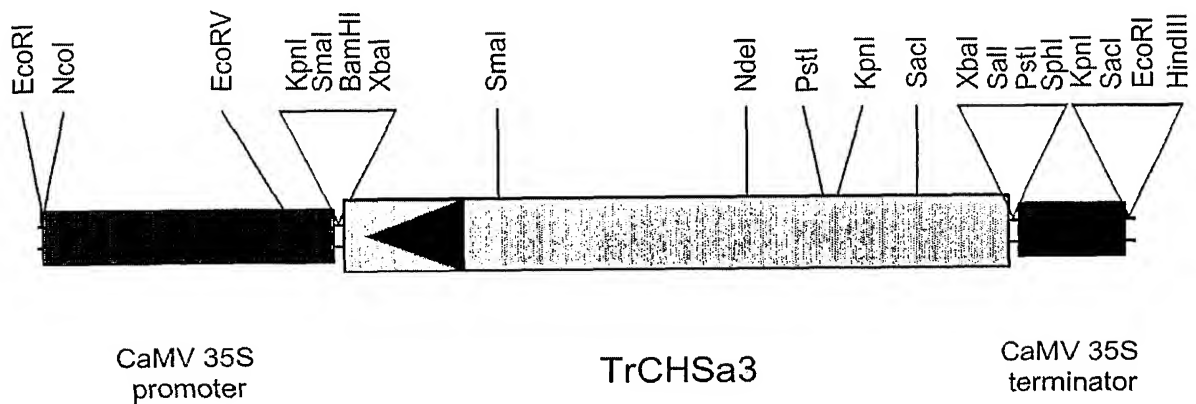
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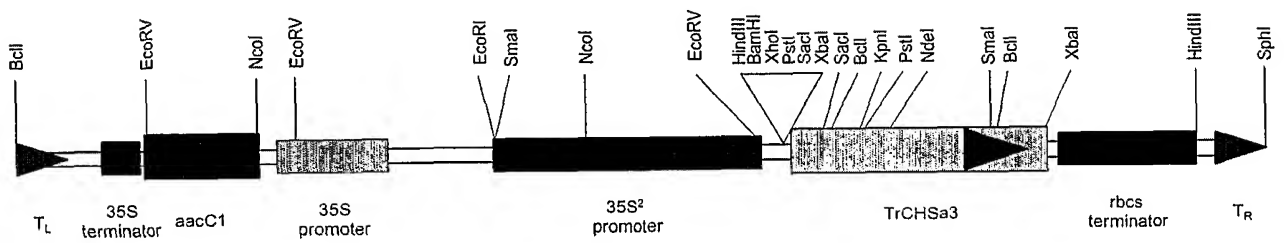
## FIGURE 142

# 216/271

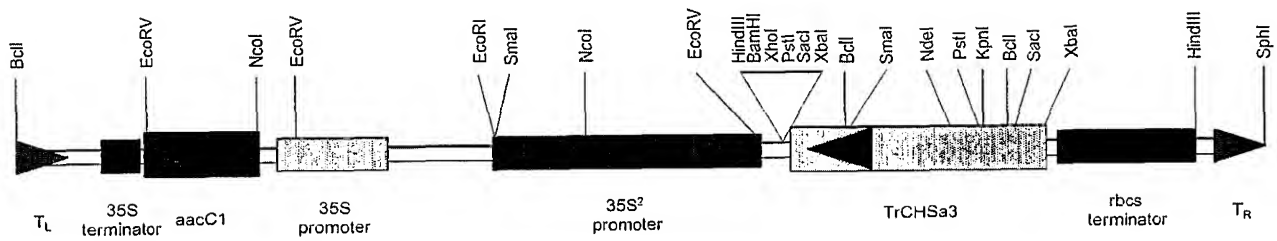
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51 ELKEKFQRM C DKSMIKSRYM YLTEEILKEN PSLCEYMAPS LDARQDMVVV  
101 EVPRLGKEAA VKAIKEWGQP KSKITHLIFC TTSGVDMPGA DYQLTKLLGL  
151 RPYVKRYMMY QQGCFAGGTV LRLAKDLAEN NKGARVLVVC SEVTAVTFRG  
201 PSDTHLDSL V GQALFGDGAA ALIVGSDPVP EIEKPIFEMV WTAQTIAPDS  
251 EGAIDGHLRE AGLTFHLLKD VPGIVSKNIN KALVEAFQPL GISDYN SIFW  
301 IAHPGGPAIL DQVEQKLALK PEKMRATREV LSEYGNMSSA CVLFILDEM R  
351 KKSAQNG LKT TGEGLDWGVL FGFGPGLTIE TVVLR SVAI

## FIGURE 143

**217/271****pDH51TrCHSa3 sense****pDH51TrCHSa3 anti****FIGURE 144**

**218/271**

pPZP221:35S²TrCHSa3 sense



pPZP221:35S²TrCHSa3 anti

**FIGURE 145**



**219/271****TrCHSc****FIGURE 146**

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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GATTCAATCT
51  GTTGTGCATA AAATTCACTC ATTGCATAGA AAACCATACA CATTTGATCT
101 TGCAAAGAAG AAATATGGGA GACGAAGGTA TAGTGAGAGG TGTCACAAAG
151 CAGACAACCC CTGGGAAGGC TACTATATTG GCTCTTGGCA AGGCATTCCC
201 TCACCAACTT GTGATGCAAG AGTGTTTAGT TGATGGTTAT TTTAGGGACA
251 CTAATTGTGA CAATCCTGAA CTTAAGCAGA AACTTGCTAG ACTTTGTAAG
301 ACAACCACGG TAAAAACAAG GTATGTTGTT ATGAATGAGG AGATACTAAA
351 GAAATATCCA GAACTTGTTG TCGAAGGCGC CTCAACTGTA AAACAACGTT
401 TAGAGATATG TAATGAGGCA GTAACACAAA TGGCAATTGA AGCTTCCCAA
451 GTTTGCCTAA AGAATTGGGG TAGATCCTTA TCGGACATAA CTCATGTGGT
501 TTATGTTTCA TCTAGTGAAG CTAGATTACC CGGTGGTGAC CTATACTTGT
551 CAAAAGGACT AGGACTAAAC CCTAAAATTC AAAGAACCAT GCTCTATTTT
601 TCTGGATGCT CGGGAGGCGT AGCCGGCCTT CGCGTTGCGA AAGACGTAGC
651 TGAGAACAAC CCTGGAAGTA GAGTTTTGCT TGCTACTTCG GAAACTACAA
701 TTATTGGATT CAAGCCACCA AGTGTTGATA GACCTTATGA TCTTGTTGGT
751 GTGGCACTCT TTGGAGATGG TGCTGGTGCA ATGATAATTG GCTCAGACCC
801 GGTATTTGAA ACTGAGACAC CATTGTTTGA GCTGCATACT TCAGCTCAGG
851 AGTTTATACC AGACACCGAG AAGAAAATTG ATGGGCGGCT GACGGAGGAG
901 GGCATAAGTT TCACACTAGC AAGGGAACCT CCGCAGATAA TCGAAGACAA
951 TGTTGAGGGA TTCTGTAATA AACTAATTGA TGTGTTGGG TTGGAGAATA
1001 AGGAGTACAA TAAGTTGTTT TGGGCTGTGC ATCCAGGTGG GCCTGCGATA
1051 TTGAATCGCG TGGAGAAGCG GCTTGAGTTG TCGCCGCAGA AGCTGAATGC
1101 TAGTAGAAAA GCTCTAATGG ATTATGGAAA TGCTAGCAGC AATACTATTG
1151 TTTATGTGCT GGAATATATG CTAGAAGAGG AAAAGAAGAT TAAAAAGGCG
1201 GGTGGAGGAG ATTCTGAATG GGGATTGATA CTTGCTTTTG GACCTGGAAT
1251 TACTTTTGAG GGGATTCTAG CAAGGAACTT GTGTGCATGA AGTCTTATAC
1301 AATTGTGATG CATGACTTAT ACTCTTATTT CTACTAATTA TTATATTAAG
1351 CAAATTCAGA ACTTTTAAAGT AATGATTTAA TGAAGAATAC TTATAGTATA
1401 TTGACTTTAT TCACTTTCAA AGCAAGTTTA TGATCCTAAG ACATGGTAGA
1451 ACTTGAGCAT GTGGAATAGT TGTAACAAAA ACTCTAAGCA AATAGAGACT
1501 TTATGTAGTA TAAAGCATTT CCAGACATGA TAAATAATGG TACCTCAGAA
1551 CATAAAATAT ATTTAGCTAT CTTTCATCCC CAACTTTACA CATCCACCAA
1601 GGTACAGAAT AAGCATATGT CAACACAAAA TGTACTCTAA GTCTAACATG
1651 AGTAACCAAA CATGATGCCT GATTAAGTTA AAAGAAAAGA AAATCTGAGG
1701 GCATAGATCT TCAATCACAC CACTCCAGAG GGAAGGCGTA GAACAAGCTG
1751 TCCGCCGAAA ACACTGCAAT TCAATAAATA TCATTAGGAC AACAGTGCAG
1801 AGTCATGCGG GAAATGTCTT AAGTCACTGT ACTAAAAATA TAGGATTATA
1851 TTATGAACTA TACTAACCTT TTCACATAAT AGTAACAGAA ATCAGCTAAG
1901 ATGAATGTCT GGACAATTTC TGAGATAAGA ACCATGACGG CCATAAGCCA
1951 TACCCCAAGG CAACCAATAA ATGTCCACGG GTATCTAACA CCTGTTGCAA
2001 GAAATAGTAA GTTATTAGGA GATGTGCGGT TACGAAATTC AAGCTACACA
2051 ACAAAGGAG GCCAGAACAA CAGCAATCTT GTAACCAGAT GACAACAATA
2101 AAATGTAAAC TTAAAGAGAC CGAACACACA AACATTGCAA CTCAGATGGA
2151 ATTGCTGCCA TGTAAGTAGT AGGAGATTTG GGACGTCAA TCAGTATATT
2201 ATGCAAATAC AAGGTATGAC CGCCTTGTCT ATTGTAGCAT ACAACAAACG
2251 TACAGTGGGT TTGTCCCTCT CAAAATGGCA GGATCTTTAC AGCACAATAT
2301 TTGGTTTTGT CATACTTATA CCATAAAAAA AAAAAAAAAA AAAAAAAAAA
2351 AAAGTACTCT GCGTTGTTAC CACTGCTTAA TCACTAGTGA ATTC

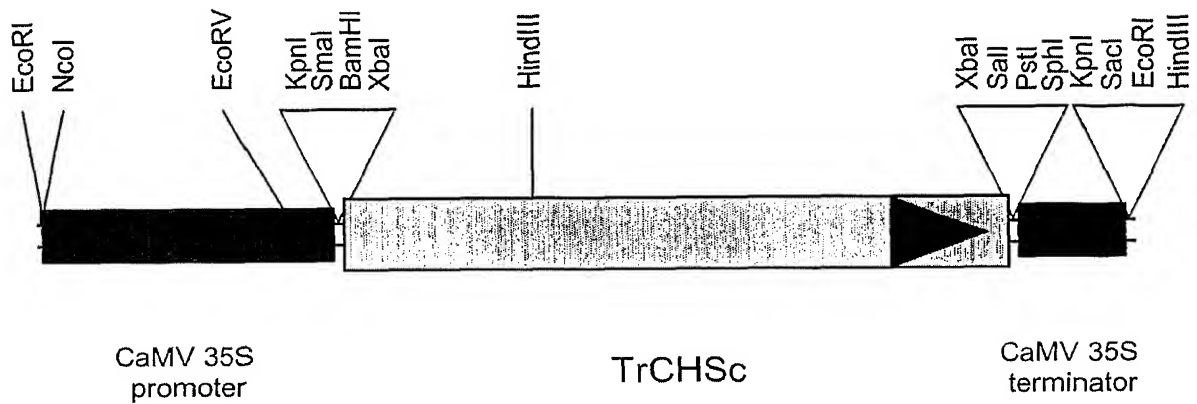
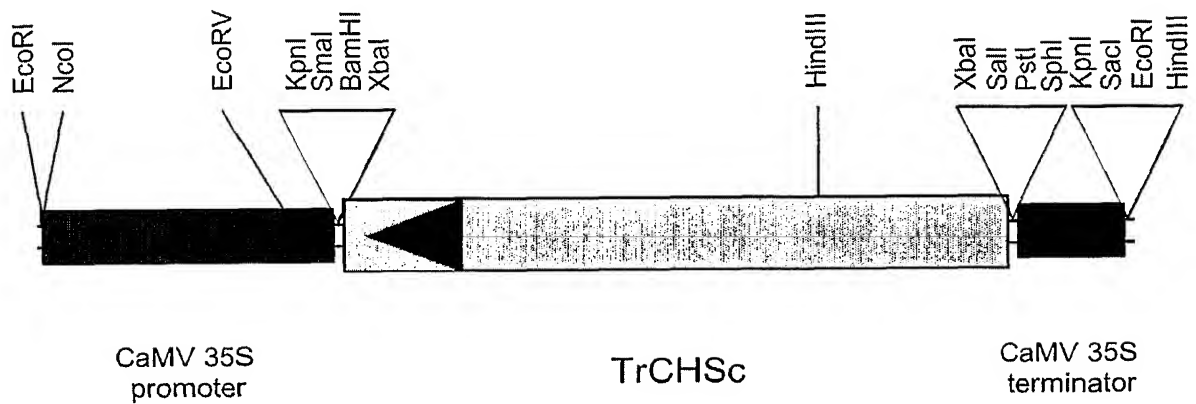
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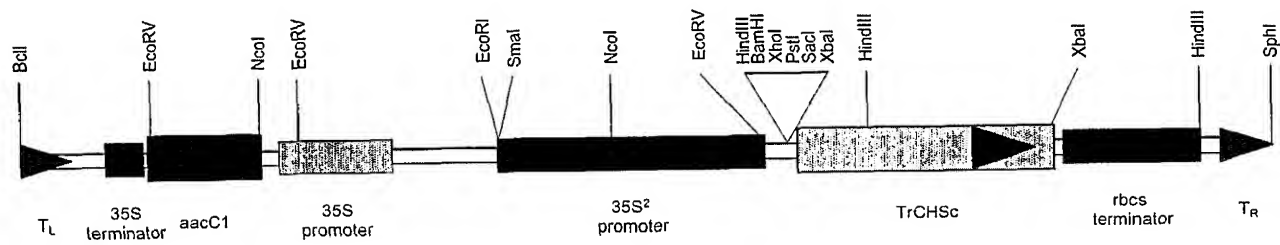
FIGURE 147

# 221/271

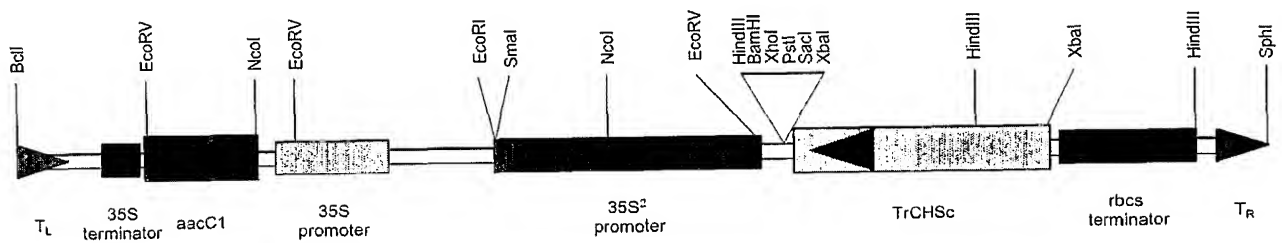
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51 PELKQKLARL CKTTTVKTRY VVMNEEILKK YPELVVEGAS TVKQRLEICN  
101 EAVTQMAIEA SQVCLKNWGR SLSDITHVVY VSSSEARLPG GDLYLSKGLG  
151 LNPKIQRMTL YFSGCSGGVA GLRVAKDVAE NNPGSRVLLA TSETTIIGFK  
201 PPSVDRPYDL VGVALFGDGA GAMIIGSDPV FETETPLFEL HTSAQEFIPD  
251 TEKKIDGRLT EEGISFTLAR ELPQIIEDNV EGFCNKLIDV VGLENKEYNK  
301 LFWAVHPGGP AILNRVEKRL ELSPQKLNAS RKALMDYGNA SSNTIVYVLE  
351 YMLEEEKKIK KAGGSDSEWG LILAFGPGIT FEGILARNLC A

## FIGURE 148

**222/271****pDH51TrCHSc sense****pDH51TrCHSc anti****FIGURE 149**

**223/271**

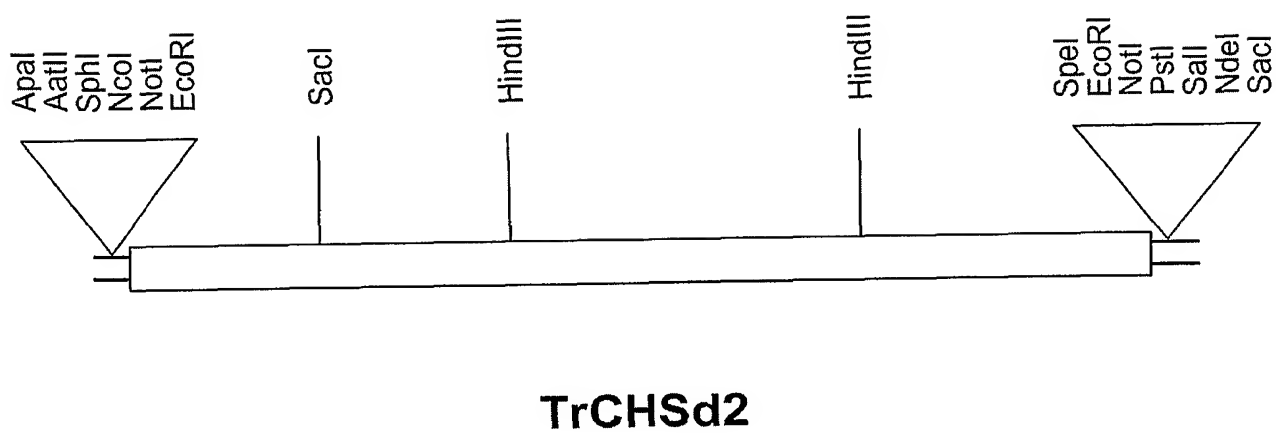
pPZP221:35S²TrCHSc sense



pPZP221:35S²TrCHSc anti

**FIGURE 150**

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**FIGURE 151**

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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GATAGCAACA
51 CACACTTTGA TTTCTTTTGT AGTCCTTGCT ACGTGGCTTT ACCAAAAAAC
101 GTTGCTAAGT CATCAACCAT TCCAATTCCT TAATATAACC TATCAGTACT
151 CACCATCTTT TCTTCCTCCC TGCTAACTTT AGACTCAGAG AAGATGGTGA
201 ATGTTAATGA GATCCGCCAG GCACAGAGAG CTGAAGGCCC TGCCACCGTG
251 TTGGCAATCG GCACTGCAAC TCCTCCAAAC TGTGTCGATC AGAGTACATA
301 CCCAGACTAC TACTTCCGCA TCACAAACAG TGAGCACAAG ACAGAGCTCA
351 AAGAAAAATT CCAGCGCATG TGTGACAAAT CTATGATTAA GAAGAGATAC
401 ATGCATTTGA CAGAAGAGAT TTTGAAGGAG AATCCAAGTT TATGTGAGTA
451 CATGGCACCT TCATTGGATG CAAGACAAGA CATGGTGGTT GTGGAAGTAC
501 CAAGGCTAGG AAAAGAGGCT GCAACAAAGG CTATCAAGGA ATGGGGTCAA
551 CCTAAGTCCA AGATTACTCA CCTCATCTTT TGCACCACAA GTGGTGTGGA
601 CATGCCTGGC GCCGACTATC AGCTTACAAA GCTTTTAGGC CTTCGTCCGC
651 ATGTGAAGCG TTATATGATG TACCAACAAG GTTGTTTCGC TGGTGGTACG
701 GTGCTTCGTT TGGCTAAAGA CTTGGCTGAA AACAACAAAG GTGCCCCGTGT
751 GTTGGTGGTT TGTTCAGAGA TCACTGCGGT TACTTCCGT GGACCCAGTG
801 AACTCATCTT TGATAGCCTT GTGGGGCAAG CATTGTTTGG AGATGGTGCA
851 GCAGCTGTGA TTGTAGGTTT AGACCCATTA CCACAAGTTG AGAAGCCCTT
901 GTTTGAATTG GTATGGACTG CTCAAACAAT CCTTCCAGAC AGTGAAGGAG
951 CCATTGATGG GCACCTTCGT GAAGTCGGGC TGACATTCCA TCTCCTCAAG
1001 GATGTTTCCTG GACTCATCTC AAAGAACATT GAGAAAGCTC TTGTTGAGGC
1051 CTTTCAACCT TTAGGTATCT CTGATTACAA TTCTATATTT TGGATCGCAC
1101 ATCCTGGTGG ACCTGCAATT CTGGACCAAG TGGAAGCCAA ATTAAGCTTA
1151 AAGCCAGAGA AAATGCAAGC CACCCGGCAT GTGCTTAGCG AGTATGGTAA
1201 CATGTCAAGT GCATGTGTGT TATTTATCTT GGATGAGATG AGGAGGAAGT
1251 CAAAAGAAGA TGGACTTGCC ACAACAGGCG AGGGGCTGGA ATGGGGTGTA
1301 CTATTTCGGTT TTGGACCCGG ACTCACTGTT GAGACTGTAT TGCTCCATAG
1351 TGTTGCCACT TAAATTGCCT AGATATGCTA TAACTATATG CTTATTTAAT
1401 TCTTTGTTTC TGGGGGATTT TATCTTCACT TACTTCACTG AGCATTTGAA
1451 TAAAGTTTGT TTTAATTATT CATAATGTAA TATGGTGTG CTTAATGTAC
1501 CCATCCATAT AATATTTGTA ATACATATAT TAATCAACTT GCAATTTTCAT
1551 GAAAAAAAAA AAAAAAAAAA AAAAAAAAAA AAAAAAAAAA AAGGAAAAAA
1601 AAAAAAAAAA AAAAAAAAAA AAGTACTCTG CGTTGTTACC ACTGCTTAAT
1651 CACTAGTGAA TTC

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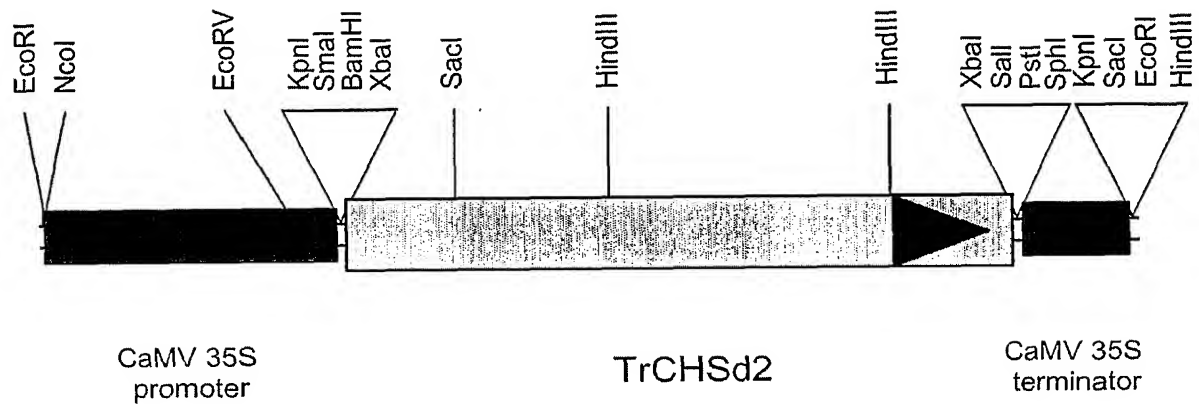
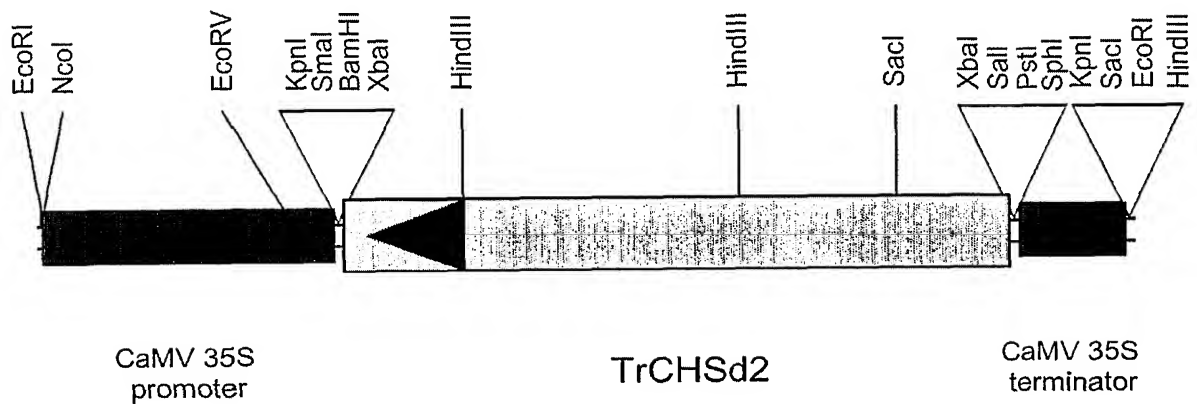
## FIGURE 152

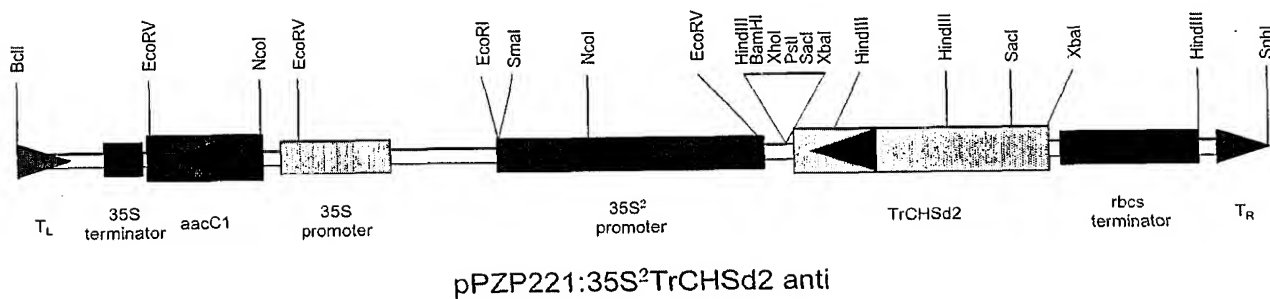
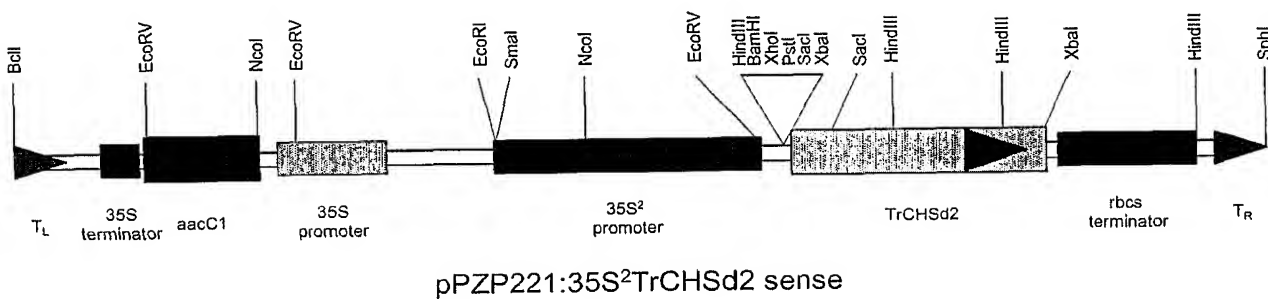
# 226/271

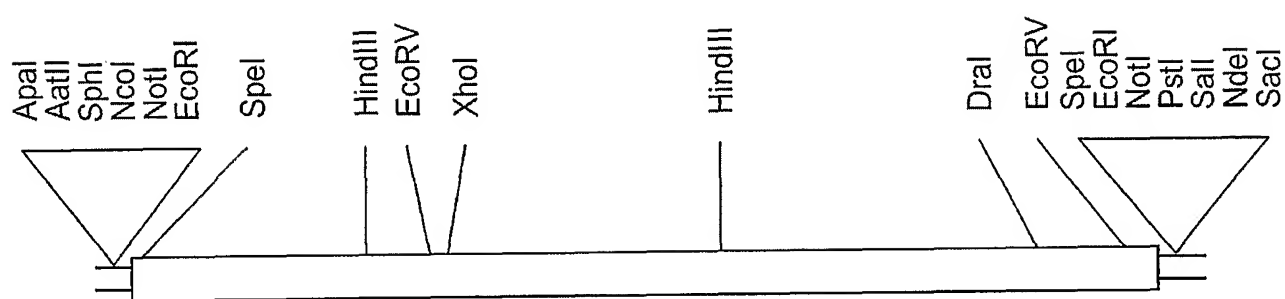
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51 ELKEKFQRM C DKSMIKKRYM HLTEEILKEN PSLCEYMAPS LDARQDMVVV  
101 EVPRLGKEAA TKAIKEWGQP KSKITHLIFC TTSGVDMPGA DYQLTKLLGL  
151 RPHVKRYMMY QQGCFAGGTV LRLAKDLAEN NKGARVLVVC SEITAVTFRG  
201 PSDTHLDSL V GQALFGDGAA AVIVGSDPLP QVEKPLFELV WTAQTILPDS  
251 EGAIDGHLRE VGLTFHLLKD VPGLISK NIE KALVEAFQPL GISDYN SIFW  
301 IAHPPGGPAIL DQVEAKLSLK PEKMQATRHV LSEYGNMSSA CVLFILDEMR  
351 RKSKE DGLAT TGEGLEWGV L FGFGPGLTVE TVLLHSVAT

## FIGURE 153



**227/271****pDH51TrCHSd2 sense****pDH51TrCHSd2 anti****FIGURE 154**

**228/271****FIGURE 155**

**229/271****TrCHSf****FIGURE 156**

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```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG ACTAAGCCTT
51 GATTCATTGT TTGTTTCCAT AACACAAGAA CTAGTGTTTG CTTGAATCTT
101 AAGAAAAAAT GCCTCAAGGT GATTTGAATG GAAGTTCCTC GGTGAATGGA
151 GCACGTGCTA GACGTGCTCC TACTCAGGGA AAGGCAACGA TACTTGCATT
201 AGGAAAGGCT TTCCCCGCCC AGGTCCTCCC TCAAGAGTGC TTGGTGGAAG
251 GATTCATTTCG CGACACTAAG TGTGACGATA CTTATATTAA GGAGAAATTG
301 GAGCGTCTTT GCAAAAACAC AACTGTGAAA ACAAGATACA CAGTAATGTC
351 AAAGGAGATC TTAGACAAC TCCAGAGCT AGCCATAGAT GGAACACCAA
401 CAATAAGGCA AAAGCTTGAA ATAGCAAATC CAGCAGTAGT TGAAATGGCA
451 ACAAGAGCAA GCAAAGATTG CATCAAAGAA TGGGGAAGGT CACCTCAAGA
501 TATCACACAC ATAGTCTATG TTTCTCGAG CGAAATTCGT CTACCCGGTG
551 GTGACCTTTA TCTTGCAAAT GAATCGGCT TAAACAGCGA TGTTAATCGC
601 GTAATGCTCT ATTTCTCTCG TTGCTACGGC GGTGTCACTG GCTTACGTGT
651 CGCCAAAGAC ATCGCCGAAA ATAACCCTGG TAGTAGGGTG TTACTCACAA
701 CATCCGAGAC CACTATTCTC GGTTTTCGAC CACCGAGTAA AGCTAGACCT
751 TATGACCTCG TTGGCGCTGC ACTTTTCGGT GATGGCGCCG CTGCTGCAAT
801 AATTGGAACA GACCCTATAT TGAATCAAGA ATCACCTTTC ATGGAATTGA
851 ACCATGCAGT CCAAAAATTC TTGCCTGATA CACAAAATGT GATTGATGGT
901 AGAATCACTG AAGAGGGTAT TAATTTTAAG CTTGGAAGAG ACCTTCCTCA
951 AAAAATTGAA GACAATATTG AAGAATTTTG CAAGAAAATT ATGGCTAAAA
1001 GTGATGTTAA GGAATTTAAT GACTTATTTT GGGCTGTTCA TCCTGGTGGG
1051 CCAGCTATAC TCAATAAGCT AGAAAATATA CTCAAATTGA AAAGTGATAA
1101 ATTGGATTGT AGTAGGAAGG CATTAATGGA TTATGGAAAT GTTAGTAGCA
1151 ATACTATATT CTATGTGATG GAGTATATGA GAGATTATTT GAAGGAAGAT
1201 GGAAGTGAAG AATGGGGATT AGGATTGGCT TTTGGACCAG GGATTACTTT
1251 TGAAGGGGTT CTCCTCCGTA GCCTTTAATC TTGAAATAAT AATTCATATG
1301 AAATTACTTG TCTTAAGATT GTGATAGGAA GATGAATATG TATTGGATTA
1351 ATATTGATAT GGTGTTATTT TAAGTTGATT TAAAAAAAAG TTTATTAATA
1401 AAGTATGATG TAACAATTGT TGTTTGAATG TAAAAGGGA AGTATACTAT
1451 TTTAAGTTCT TGACCATACT GATTTTTTCT TTACACATTT TCATATCTAA
1501 AATTGTTCTA TGATATCTTC ATTGTTGATA CTGTAATAAT ATAATATCTA
1551 ATTTGGCTGG CAAAATGAAA GATTTTTCAC CGAAAAAAA AAAAAAAAAA
1601 AAAAAAAAAA AAGTACTCTG CGTTGTTACC ACTGCTTAAT CACTAGTGAA
1651 TTC

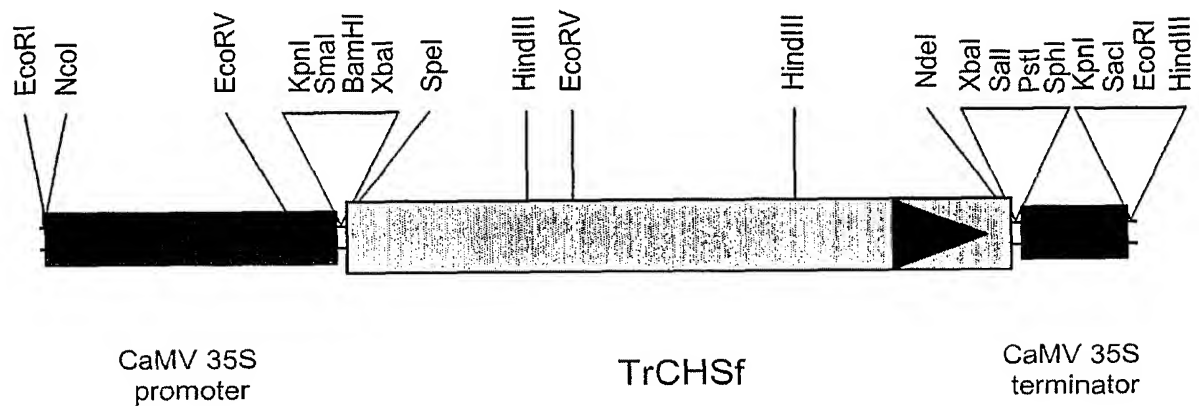
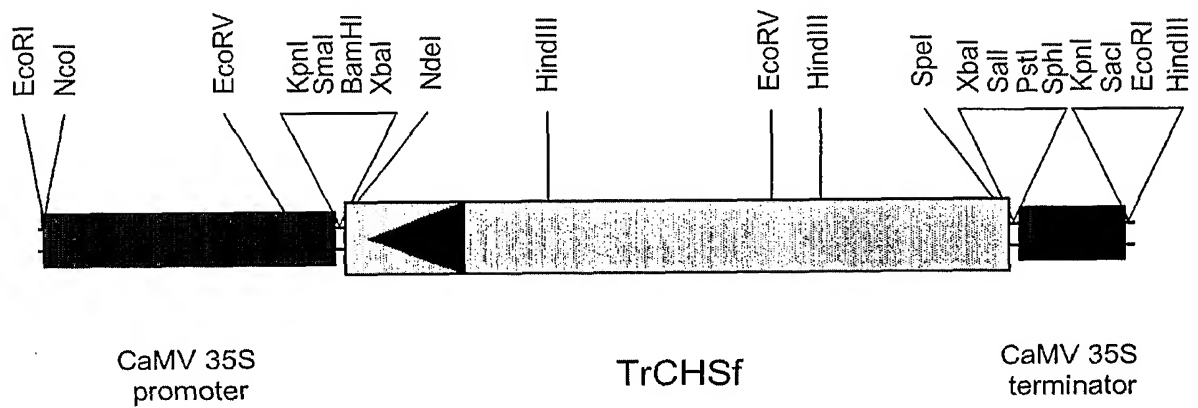
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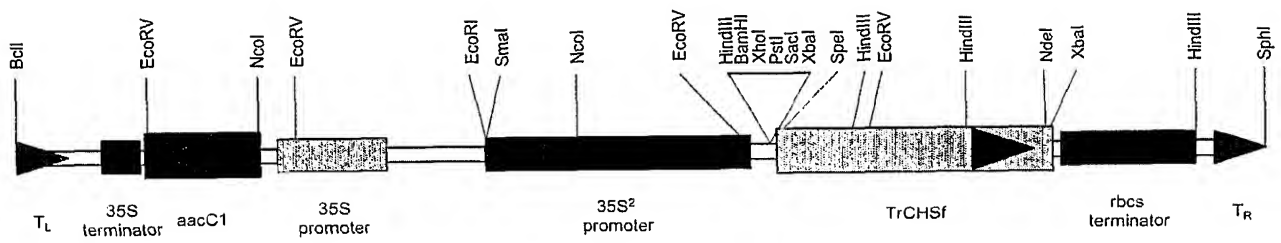
## FIGURE 157

# 231/271

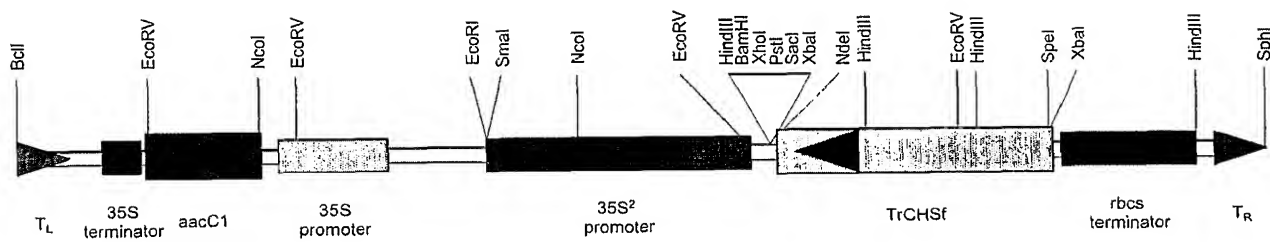
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1  MPQGDNLNGSS  SVNGARARRA  PTQGKATILA  LGKAFPAQVL  PQECLVEGFI
51  RDTKCDDTYI  KEKLERLCKN  TTVKTRYTVM  SKEILDNYPE  LAIDGTPTIR
101 QKLEIANPAV  VEMATRASKD  CIKEWGRSPQ  DITHIVYVSS  SEIRLPGGDL
151 YLANELGLNS  DVNRVMLYFL  GCYGGVTGLR  VAKDIAENNP  GSRVLLTTSE
201 TTILGFRPPS  KARPYDLVGA  ALFGDGAAAA  IIGTDPILNQ  ESPFMELNHA
251 VQKFLPDTQN  VIDGRITEEG  INFKLGRDLP  QKIEDNIEEF  CKKIMAKSDV
301 KEFNDLFWAV  HPGGPAILNK  LENILKLKSD  KLDCSRKALM  DYGNVSSNTI
351 FYVMEYMRDY  LKEDGSEEWG  LGLAFGPGIT  FEGVLLRSL
```

## FIGURE 158

**232/271****pDH51TrCHSf sense****pDH51TrCHSf anti****FIGURE 159**

**233/271**

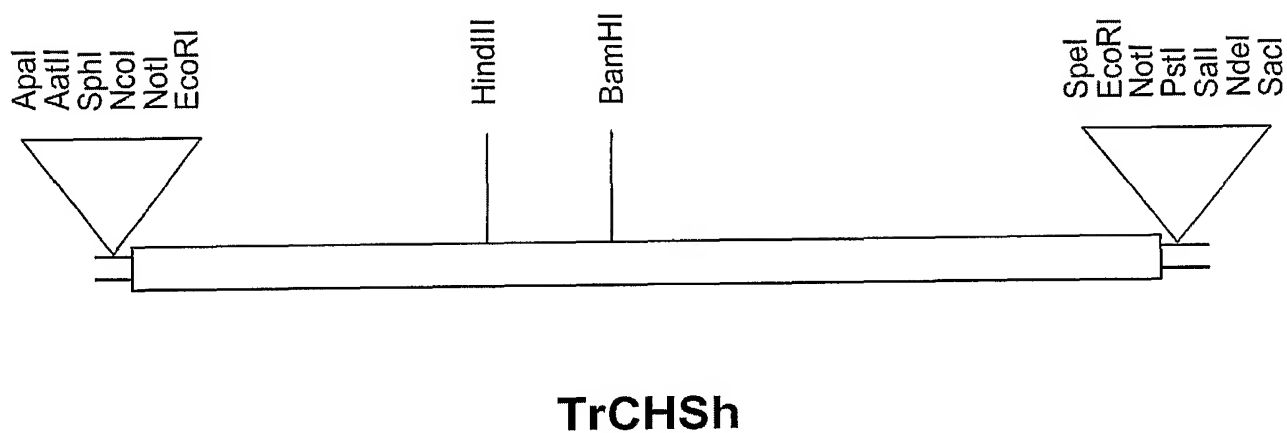
pPZP221:35S<sup>2</sup>TrCHSf sense



pPZP221:35S<sup>2</sup>TrCHSf anti

**FIGURE 160**

**234/271**



**FIGURE 161**



# 235/271

```

1  GAATTCACTA GTGATTAAGC AGTGGTAACA ACGCAGAGTA CGCGGGGGGAA
51 TCCACCAAAT CAACACCATT AATAACCTTC CAAATTCTCG TTACCTCACC
101 AAATCTCATT TTTCATTATA TATCTTGGGT ACATCTTTTG TTACCTCCAA
151 CAAAAAATG GTGACCGTAG AAGAGATTTCG TAACGCCCAA CGTTCAAATG
201 GCCCTGCCAC TATCTTAGCT TTTGGCACAG CCACTCCTTC TAACTGTGTC
251 ACTCAAGCTG ATTATCCTGA TTACTACTTT CGTATCACCA ACAGCGAACA
301 TATGACTGAT CTTAAGGAAA AATTCAAGCG GATGTGTGAT AGATCAATGA
351 TAAAGAAACG TTACATGCAC CTAACAGAAG ACTTTCTGAA GGAGAATCCA
401 AATATGTGTG AATACATGGC ACCATCACTA GATGTAAGAC GAGACATAGT
451 GGTGTGTGAA GTACCAAAGC TAGGTAAAGA AGCAGCAAAA AAAGCCATAT
501 GTGAATGGGG ACAACCAAAA TCCAAAATCA CACATCTTGT TTTCTGCACC
551 ACTTCCGGTG TTGACATGCC GGGAGCCGAT TACCAACTCA CCAAACCTTTT
601 AGGCTTAAAA CCTTCTGTCA AGCGTCTCAT GATGTATCAA CAAGGTTGTT
651 TCGCTGGCGG CACAGTTCTC CGCTTAGCAA AAGACCTTGT TGAGAATAAC
701 AAAAATGCAA GAGTTCTTGT TGTTTGTTCT GAAATTACTG CGGTACTTTT
751 TCGTGGACCA TCGGATACTC ATCTTGATTG GCTCGTGGGA CAGGCGCTTT
801 TTGGTGATGG AGCCGCAGCA ATGATTATTG GTGCGGATCC TGATTTAACC
851 GTGGAGCGTC CGATTTTCGA GATTGTTTCG GCTGCTCAGA CTATCTTCC
901 TGATTCTGAT GGC GCAATTG ATGGACATCT TCGTGAAGTG GGGCTCACTT
951 TTCATTTATT GAAAGATGTT CCGGGGATTA TTTCAAAGAA CATTGAAAAA
1001 AGTTTAGTTG AAGCTTTTGC GCCTATTGGG ATTAATGATT GGAAC TCAAT
1051 ATTTTGGGTT GCACATCCAG GTGGACCGGC TATTTTAGAC CAGGTTGAAG
1101 AGAAACTCCA TCTTAAAGAG GAGAAACTCC GGTCCACCCG GCATGTGCTT
1151 AGTGAATATG GAAATATGTC AAGTGCATGT GTTTTATTTA TTTTGGATGA
1201 AATGAGAAAG AGGTCTAAAG AGGAAGGGAT GATTACAAC T GGTGAAGGGT
1251 TGGAATGGGG TGTGTTGTTT GGGTTTGGAC CGGGTTTAA C TGTTGAAACC
1301 GTTGTGCTTC ATAGTGTTCC GGTT CAGGGT TGAATTTATT ATACATAGAT
1351 TGGA AAAATAA AATTTGCCTG CCGAGAGATG TGA ACTAACT TTGTAGGCAA
1401 GCTCAAATTA AAGTTTGAGA TAATATTGTG CTTTAGTTAT TATGGTATGT
1451 AATGTAATGT TTTTACTTTT TTCGAAATTC ATGTAATTTG ATATGTAAAG
1501 TAATATGTTT GGGTTGGAAT ATAATTATTT GTTAACTAAA AAAAAAAAAA
1551 AAAAAAAAAA AAAAAGTACT CTGCGTTGTT ACCACTGCTT AATCGAATTC

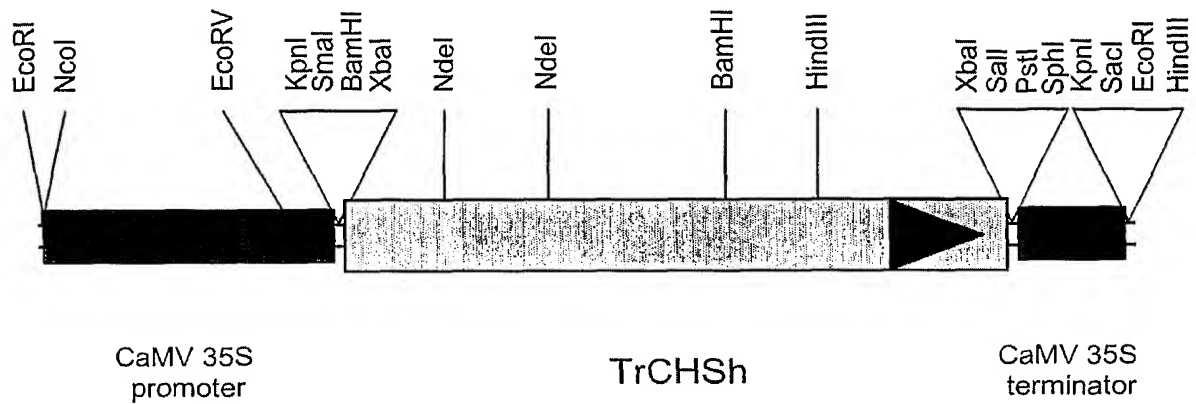
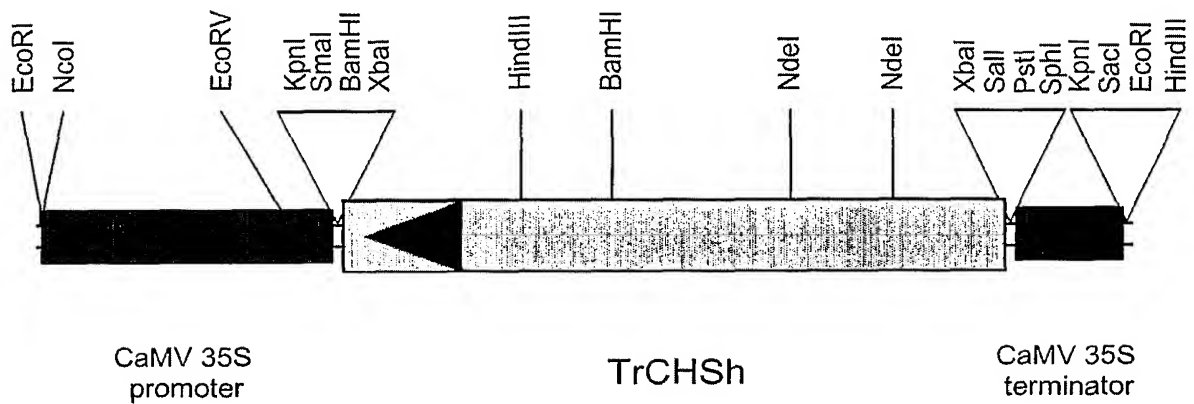
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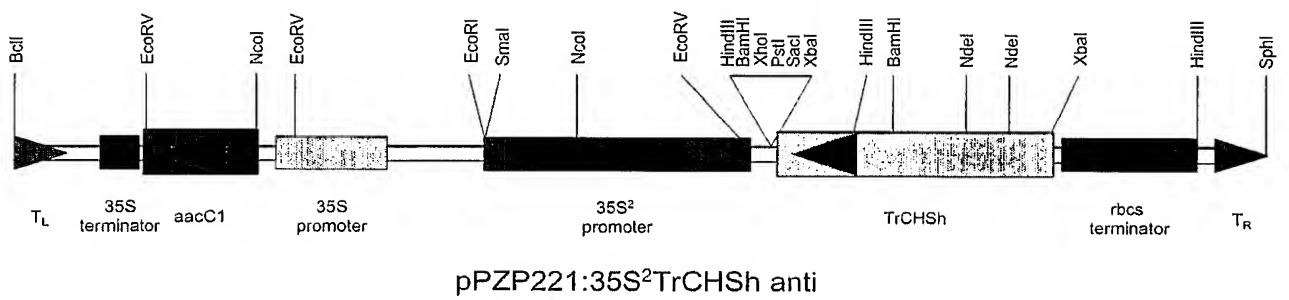
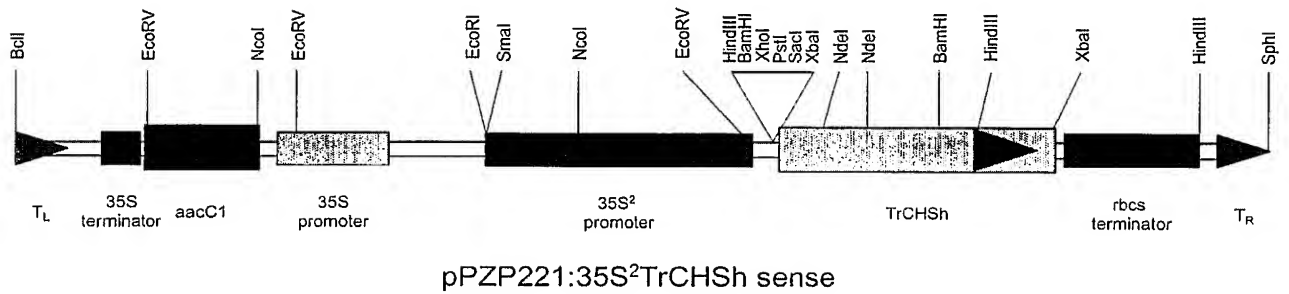
## FIGURE 162

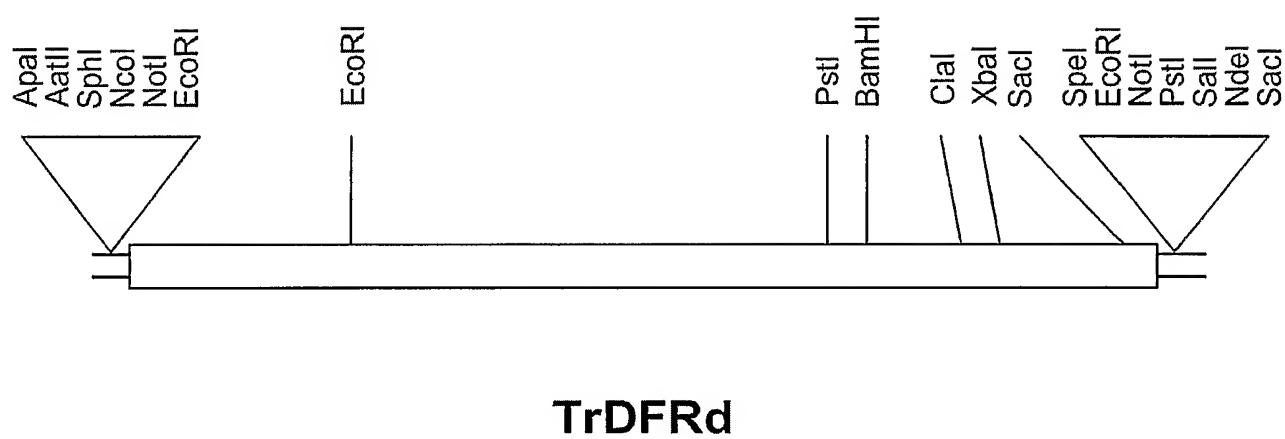
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1 MVTVEEIRNA QRSNGPATIL AFGTATPSNC VTQADYPDYY FRITNSEHMT  
51 DLKEKFKRMC DRSMIKKRYM HLTEDFLKEN PNMCEYMAPS LDVRRDIVVV  
101 EVPKLGKEAA KKAICEWGQP KSKITHLVFC TTSGVDMPGA DYQLTKLLGL  
151 KPSVKRLMMY QQGCFAGGTV LRLAKDLVEN NKNARVLVVC SEITAVTFRG  
201 PSDTHLDSLV GQALFGDGAA AMIIGADPDL TVERPIFEIV SAAQTILPDS  
251 DGAIDGHLRE VGLTFHLLKD VPGIISKNIE KSLVEAFAPI GINDWNSIFW  
301 VAHPGGPAIL DQVEEKLHLK EEKLRSTRHV LSEYGNMSSA CVLFILDEMR  
351 KRSKEEGMIT TGEGLEWGV LFGFGPGLTVE TVVLHSVPVQ G

## FIGURE 163

**237/271****pDH51TrCHSh sense****pDH51TrCHSh anti****FIGURE 164**

**238/271****FIGURE 165**

**239/271****FIGURE 166**

# 240/271

```

1  GAATTCACTA GTGATTAAGC AGTGGTAACA ACGCAGAGTA CGCGGGGGTG
51 ACTTGATCTA GCAGTTATCA AACACAACAG TCTTCCACTT GAGCTCTGTT
101 TCTCCACATG TCGAAGCTAG TTTGCGTCAC CGGCGGCAGC GGATGCATCG
151 GTTCATGGCT AGTCCATCTC CTTCTCCTCC GCGGCTACAC TGTTACAGCC
201 ACCGTCCAAA ATCTCAATGA TGAGAACGAA ACGAAGCATC TAGAAGCTCT
251 CGAAGGAGCA CAAACTAATC TCCGTCTCTT CCAGATCGAT CTCCTTAACT
301 ACGACACAAT CCTCGCTGCT GTCCGCGGTT GCGTCGGAAT TTTCCACCTC
351 GCTTCACCTT GCACTGTAGA CAAAGTTCAT GATCCTCAGA AGGAGCTTTT
401 GGATCCTGCA ATTAAAGGGA CTTTGAATGT GCTTACTGCA GCTAAGGAAG
451 TAGGGGTGAA GCGTGTGGTT GTTACCTCGT CTGTCTCGGC GATTACTCCT
501 AGTCCTGATT GGCCTTCTGA TGTTGTTAAA AGAGAGGATT GTTGGACTGA
551 TGTTGAATAT TGCAAGAAAA AAGAGTTGTG GTATCCGTTG TCCAAAACAT
601 TGGCTGAGAA AGCTGCGTGG GATTTTTTCCA AAGAAAATGG TTTGGATGTT
651 GTTGTGGTGA ATCCCGGTAC TGTGATGGGT CCTGTTATTC CACCACGGCA
701 TAATGCAAGC ATGCTCATGC TTGTGAGACT TCTTGAAGGC TGCGCTGAAA
751 CATTTGAAGA CTATTTTATG GGATTGGTCC ACTTCAAAGA TGTAGCATTG
801 GCGCATATTT TGGTGTATGA GAACAAAGAA GCATCTGGTA GACATGTGTG
851 TGTTGAAACT ATCTCTCACT ACGGTGATTT TGTGGCAAAA GTTGCTGAAC
901 TTTATCCAGA ATATAGTGTT CCTAGGATGC AGCGAGATAC GCAACCTGGA
951 TTGTTGAGAG CGAATGATGG ATCAAAGAAG CTCATAGATT TGGGTTTGGA
1001 ATTCATTCCA ATGGAGCAAA TTATCAAGGA TGCTGTAGAG AGTTTGAAGA
1051 ACAAAGGATT CATTTCTTGA ATGATGTTAC TGTTCTTTGG AGAACCTTAT
1101 AGTTACCAGA GTATAGACTA AATAATATAT AGGTGATGGG TCAGAGAATG
1151 AGTACTTATG TCATGAGTTG TGTCTGTATA ATATGTTTTT TCAATTCTTA
1201 TATGTTAAAT TGCTAATGTT AACTTCAATA TTTATCAGCC AGTATTGTTT
1251 TTTTAATAAA ATATTGAAGC AAAAAAAAAA AAAAAAAAAA AAAAAAAGT
1301 ACTCTGCGTT GTTACCACTG CTTAATCGAA TTC

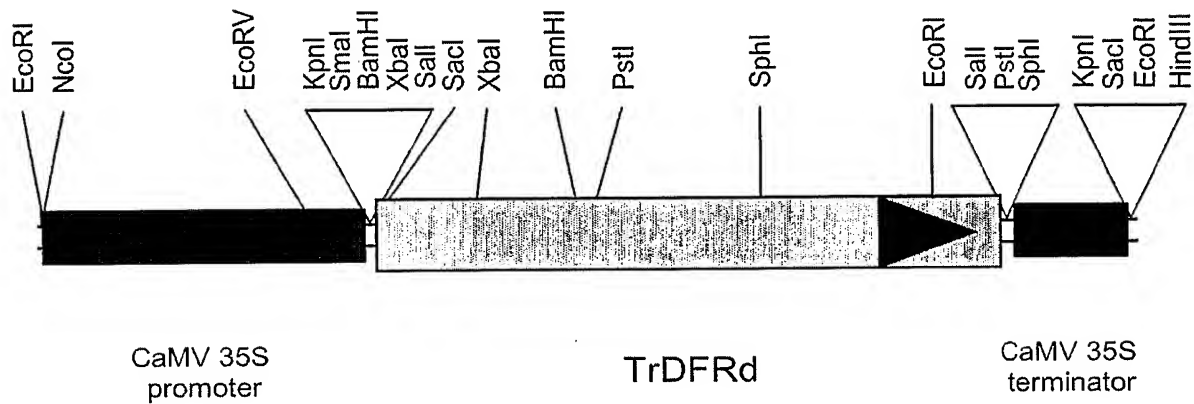
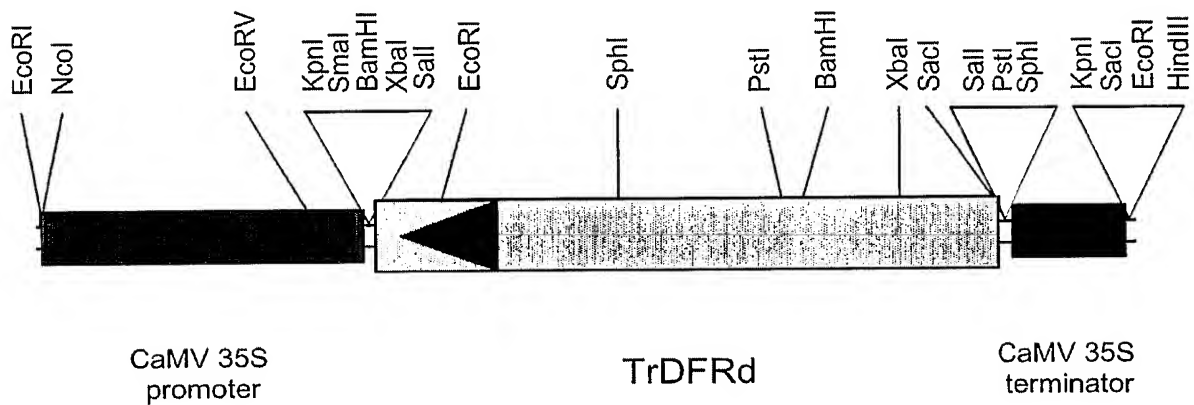
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## FIGURE 167

# 241/271

1 MSKLVCVTGG SGCIGSWLVH LLLLRGYTVH ATVQNLNDEN ETKHLEALEG  
51 AQTNLRLFQI DLLNYDTILA AVRGCVGIFH LASPCTVDKV HDPQKELLD  
101 AIKGTLNVLT AAKEVGVKRV VVTSSVSAIT PSPDWPSDVV KREDCWTDVE  
151 YCKKKELWYP LSKTLAEKAA WDFSKENGLD VVVVNP GTVM GPVIPPRHNA  
201 SMLMLVRLLE GCAETFEDYF MGLVHFKDVA LAHILVYENK EASGRHVCVE  
251 TISHYGDFVA KVAELYPEYS VPRMQRDTQP GLLRANDGSK KLIDLGLEFI  
301 PMEQUIIKDAV ESLKNKGFIS

## FIGURE 168

**242/271****pDH51TrDFRd sense****pDH51TrDFRd anti****FIGURE 169**



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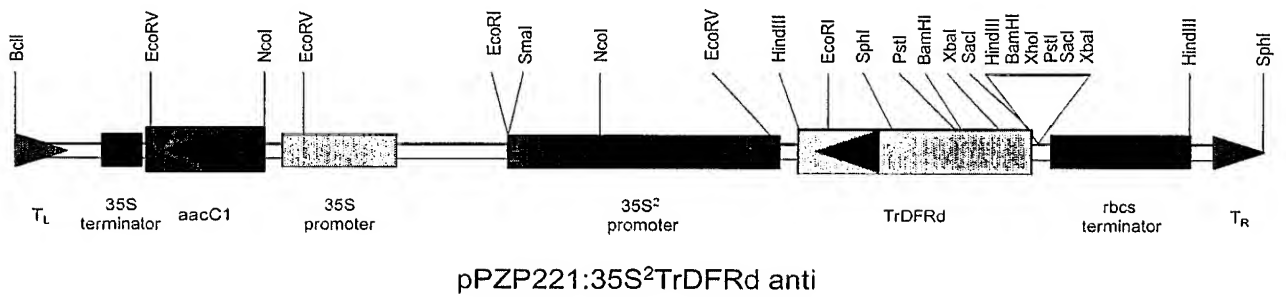
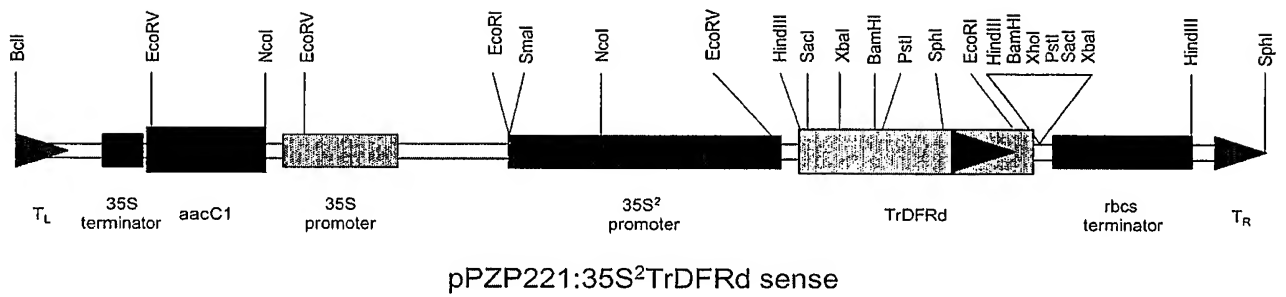
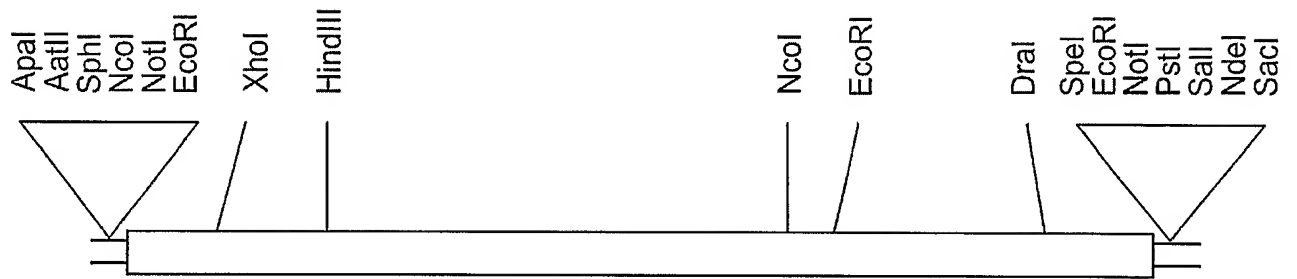


FIGURE 170

244/271



TrF3Ha

**FIGURE 171**

# 245/271

```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GACCACTCTA
51  TTTATTTCTA CTTAAACCTC ACAAAAAATA AACCACACAA CACACAAACA
101 CCAAAAACAG AGCACCGTTT CCATCATCAA ACAATGGCAC CAGCCAAAAC
151 TCTAAGTTAT CTCTCACAAC AAAACACTCT CGAGTCAAGT TTCGTTAGGG
201 AAGAAGATGA GCGTCCAAAA GTTGCCCTACA ATAACCTCAG CAACGAGATT
251 CCAATCATTT CTCTTGCTGG AATTGATGAG GTTGATGGTC GTAGAACAGA
301 GATATGTAAC AAGATTGTTG AAGCTTGTTGA GAATTGGGGT ATTTTTCAGG
351 TTGTTGATCA TGGTGTTGAT ACAAACCTTG TTTCTGAGAT GACCCGTTTT
401 GCTAGAGAGT TTTTTCCTTT GCCACCGGAA GAGAAGCTCC GGTTCGACAT
451 GTCCGGTGGT AAAAAGGGTG GTTTCATTGT CTCTAGTCAT CTTCAAGGAG
501 AAGCAGTGAA GGATTGGAGA GAGCTAGTGA CATATTTTTC ATACCCAATT
551 AAACAAAGAG ATTATTC AAG GTGGCCAGAC AAGCCAGAAG GATGGAAAGA
601 GGTAACAGAA AAATACAGTG AAAACCTAAT GAATTTAGCT TGCAAGCTAT
651 TGGAAGTTTT ATCAGAAGCA ATGGGTTTAG AAAAAGAAGC TCTAACAAAA
701 GCATGTGTTG ATATGGATCA AAAAGTTGTT ATAAATTATT ACCCAAAATG
751 CCCTGAACCT GACCTCACAC TTGGCCTTAA ACGTCACACT GACCCTGGCA
801 CAATTACTCT TTTGCTTCAA GATCAAGTTG GTGGTCTTCA AGCTACCAAA
851 GATAATGGTA AGACGTGGAT TACAGTTCAA CCAGTTGAAG GTGCTTTTGT
901 TGTTAATCTT GGAGACCATG GTCACATCT AAGTAATGGA CGGTTCAAAA
951 ATGCTGACCA TCAAGCAGTG GTGAATTCGA ACTACAGCCG TTTATCAATA
1001 GCAACATTTT AAAATCCAGC TCCAGATGCA ACTGTGTACC CTTTGAAGAT
1051 TAGAGATGGT GAAAAATCTG TGTTGGAAGA ACCAATCACT TTTGCTGAAA
1101 TGTATAGAAG GAAGATGACC AAAGACCTTG AAATTGCTAG GATGAAGAAG
1151 TTGGCTAAGG AACAACAAC TTAGGGACTTG GAGGAGAACA AGACTAAATA
1201 TGAGGCCAAA CCTTTGAATG AGATCTTTGC TTAATTAATT AGTCTTAATT
1251 TAAATAATAA ATTTTAGACT TAATTTACAT ATAATAATTT TAATTTTTTG
1301 TTCAATTAAT CTATGTTTAA TTTGTCGTTA TTGTCCACGT GTATTAAGCT
1351 GCTTGGTTGT GTGTGCCTTG GAGAATAATC AATAATATTA CATCTATGTT
1401 TAATTATAAA AAAAAAAAAA AAAAAAAAAA GTATCTGCGT TGTTACCACT
1451 GCTTAATCAC TAGTGAATTC

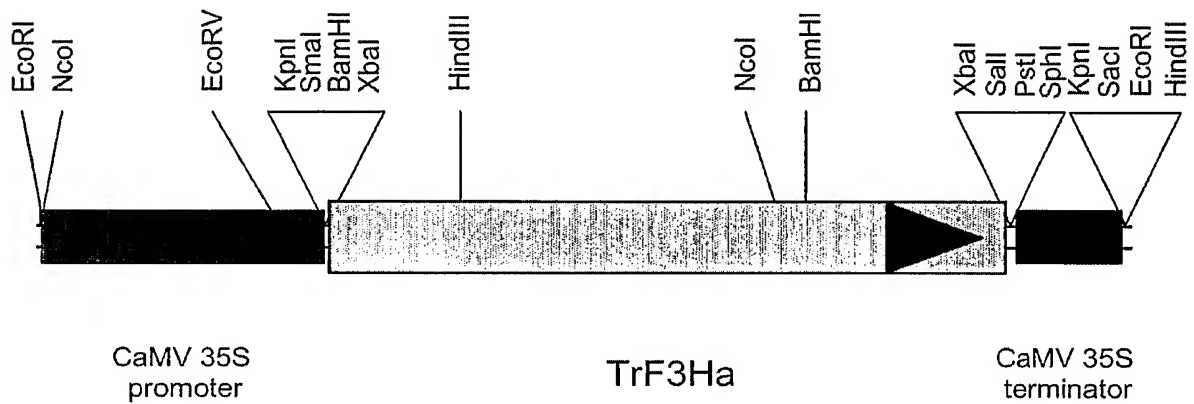
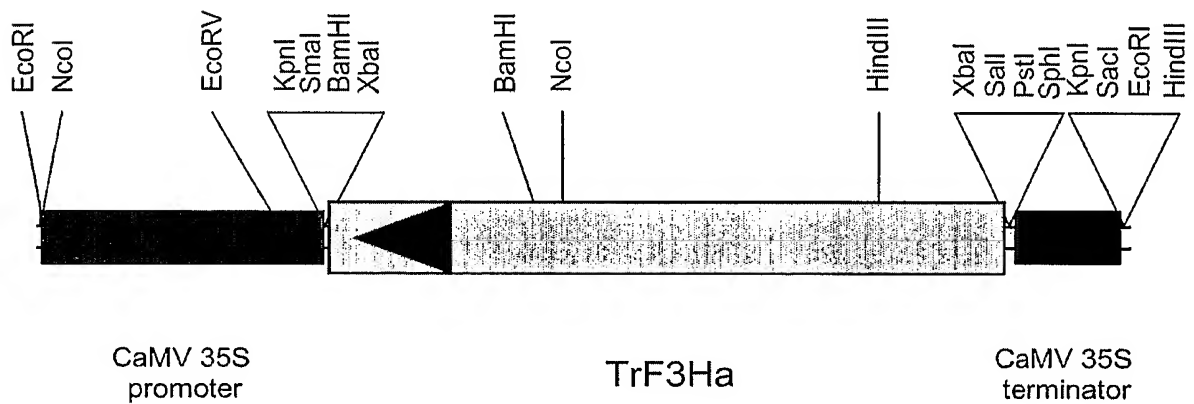
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## FIGURE 172

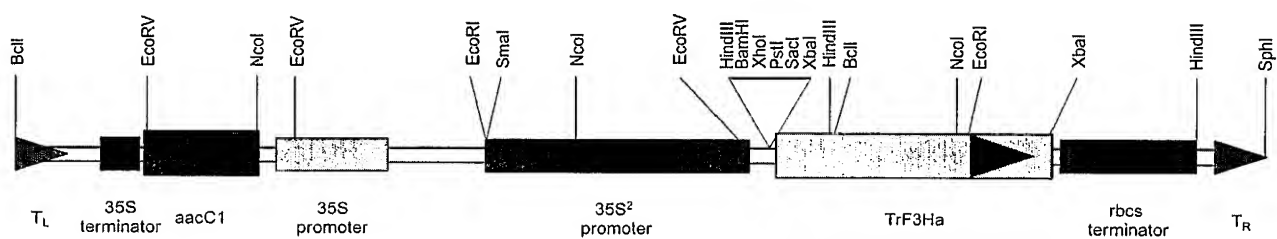
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1 MAPAKTLSYL SQQNTLESSF VREEDERPKV AYN NFSNEIP IISLAGIDEV  
51 DGR RTEICNK IVEACENWGI FQVVDHGVD T KLVSEMTRFA REFFALPPEE  
101 KLRFDMSGGK KGGFIVSSHL QGEAVKDWRE LV TYFSYPIK QRDYSRW PDK  
151 PEGWKEVTEK YSENL MNLAC KLLEVLSEAM GLEKEALTKA CVDMDQKVVI  
201 NYYPKCPEPD LTLGLKRHTD PGTITLLLQD QVGGLQATKD NGKTWITVQP  
251 VEGAFVVNLG DHGHYLSNGR FKNADHQAVV NSNYSRLSIA TFQNPAPDAT  
301 VYPLKIRDGE KSVLEEPITF AEMYRRKMTK DLEIARMKKL AKEQQLRDLE  
351 ENKTKYEAKP LNEIFA

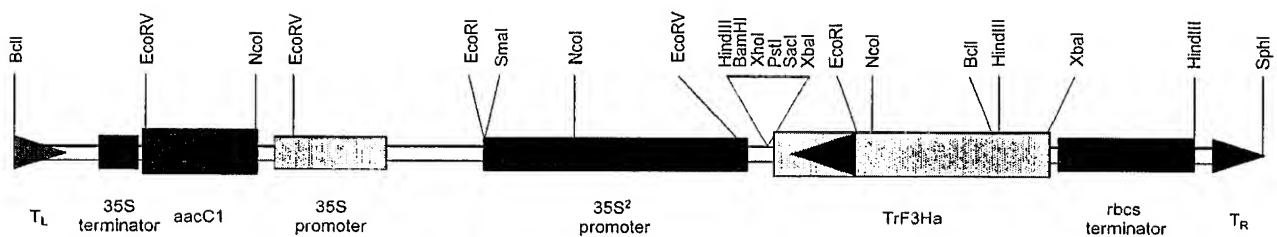
## FIGURE 173

**247/271****pDH51TrF3Ha sense****pDH51TrF3Ha anti****FIGURE 174**

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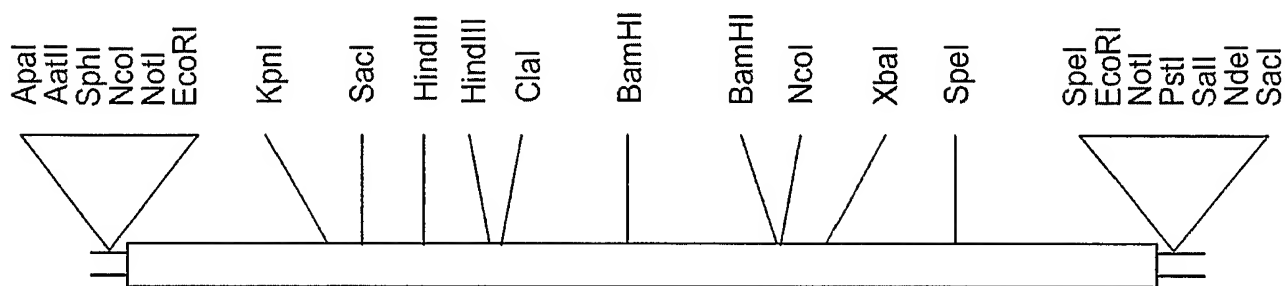


pPZP221:35S²TrF3Ha sense



pPZP221:35S²TrF3Ha anti

FIGURE 175

**249/271****TrPALa****FIGURE 176**

# 250/271

```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG GAGGAAATTC
51  ACAACTCAAA TATTCCTTTA ATTCTTTTCAT ATAATCATTT GAATTTCCAT
101 TCTCCCTAAA AATTCTATAG CTACCACATC ACACAACATA ACAAATTAAG
151 AAATATTTTAT TACTATATTA AGATATGGAA GTAGTAGCAG CAGCAATCAC
201 AAAAAACAAT GGCAAGATTG ATTCATTTTG CTTGAATCAT GCTAATGCTA
251 ATAACATGAA AGTGAATGGT GCTGATCCTT TGAATTGGGG TGTGGCTGCT
301 GAGGCAATGA AGGGAAGTCA CTTGGATGAG GTGAAGCGTA TGGTGGAGGA
351 ATACCGGAAA CCGGTTGTCC GTCTTGGTGG CGAGACACTA ACCATTTCTC
401 AGGTGGCTGC CATTGCTGCA CACGATGGTG CAACGGTGGA GCTATCGGAA
451 TCTGCTAGAG CCGGCGTTAA GGCAAGCAGT GACTGGGTTA TGGAGAGTAT
501 GAACAAAGGT ACCGACAGCT ACGGTGTCCC AACAGGGTTC GGCCTACCT
551 CGCACCGCCG AACCAAACAA GGTGGTGCTT TGCAGAAAGA GCTCATAAGG
601 TTTTTGAATG CTGGAATATT TGGAAATGGA ACTGAGTCAA GCCACACACT
651 ACCACACACA GCCACAAGAG CTGCCATGCT AGTGAGAATC AACACACTTC
701 TCCAAGGCTA TTCAGGAATT AGATTTGAAA TCTTAGAAGC TATCACCAG
751 CTTCTTAACA ACAATGTCAC CCCATGTTTA CCGCTTCGCG GTACAATCAC
801 AGCTTCAGGA GATTTAGTCC CTCTTCTTCA CATTGCTGGT TTAATAACCG
851 GACGACCAA TTCCAAGGCT CATGGACCTT CTGGAGAAGT ACTTAATGCA
901 AAACAAGCTT TTCAATCAGC TGGAATCGAT GCCGAGTTCT TTGAATTACA
951 ACCAAAAGAA GGCTTGCCC TTGTTAACGG AACCGCTGTT GGTCTGGTT
1001 TAGCTTCTAT TGTCTTTTTT GAGGCTAATA TATTGGCGGT GTTGTCTGAA
1051 GTTCTATCTG CAATTTTCGC TGAAGTTATG CAAGGGAAGC CCGAATTTAC
1101 CGATCATTTG ACACATAAGT TGAAACATCA CCCTGGTCAA ATTGAGGCTG
1151 CTGCTATTAT GGAACACATT TTGGATGGGA GTGCTTATGT TAAAGACGCT
1201 AAGAAGTTGC ATGAGATGGA TCCTTTACAG AAGCCAAAAC AAGTAGATA
1251 TGCTTCTAGG ACTTCGCCAC AATGGCTTGG TCCTTTGATT GAAGTGATTA
1301 GATTCTCTAC CAAGTCAATT GAGAGAGAGA TCAACTCTGT CAATGACAAT
1351 CCTTTGATTG ATGTTTCAAG GAACAAGGCT TTGCATGGTG GAAATTTTCA
1401 AGGAACACCT ATCGGAGTAT CCATGGATAA TACACGTTTG GCTCTTGCAT
1451 CAATTGGCAA ACTTATGTTT GCTCAATTCT CTGAGCTTGT CAATGATTTT
1501 TACAACAATG GATTGCCATC AAATCTCTCT GCTAGTAGAA ATCCGAGCTT
1551 GGATTATGGG TTCAAGGGAT CCGAAATTGC CATGGCTTCT TATTGTTCCG
1601 AGTTGCAATA TCTTGCAAAT CCGGTTACAA CTCATGTCCA AAGTGCGGAA
1651 CAACACAACC AAGATGTCAA CTCTTTGGGT TTGATTTCTT CTAGAAAAAC
1701 TTATGAAGCA ATTGAGATCC TTCAATTGAT GTCTTCCACA TTCTTGATTG
1751 CACTTTGTCA AGCAATTGAT TTAAGACATT TGGAGGAGAA TTTGAAAAAC
1801 TCGGTCAAAA ATACCGTAAG CCAAGTGGCC AAAAAGACAC TAACCATAGG
1851 TGTCAATGGA GAACTTCATC CTTCAAGATT TTGTGAAAAA GACTTATTGA
1901 AAGTGGTTGA TAGGGAACAT GTCTTTGCCT ACATTGATGA TCCTTGATAGT
1951 GCTACATACC CATTGATGCA AAAACTCAGG CAAGTACTAG TGGATCATGC
2001 ATTAGTTAAT GGAGAAAGTG AGAAGAATTT GAACACATCA ATCTTCCAAA
2051 AGATTGCAAC TTTTGAGGAA GAGTTGAAAA ACCTTTGCCA AAAGAGGTTG
2101 AAAGTGCAAG GATTGCATAT GAAAGTGGA ATTCAACAAT TCCAAACAAG
2151 ATCAATGGAT GCAGATCTTA TCCACTCTAC AATTTTGTGA GAAAGGAGTT
2201 GGAAGTGGT TTGCTAACTG GAGAAAATGT CATTTACCGG GGTGAAGAGT
2251 GTGACAACT ATTCACAGCT ATGTGTCAAG GAAAAATCAT TGATCCTCTT
2301 CTTGAATGCT TGGGAGAGTG GAACGGTGCT CCTCTTCCAA TTTGTAACT
2351 TTGATTGTTA GTTCATAAAA TGTTTTATTT GTATTTATCA TTTGTATTTA
2401 TGCGAGTGTA GTAATAATGA TTAGGTGTTT TGTGCCTTTA ATGAAAAAAA
2451 AAAAAAAAAA AAAAAAAAAA AAAAGTACTC TGCGTTGTGA CCACTGCTTA
2501 ATCACTAGTG AATTC

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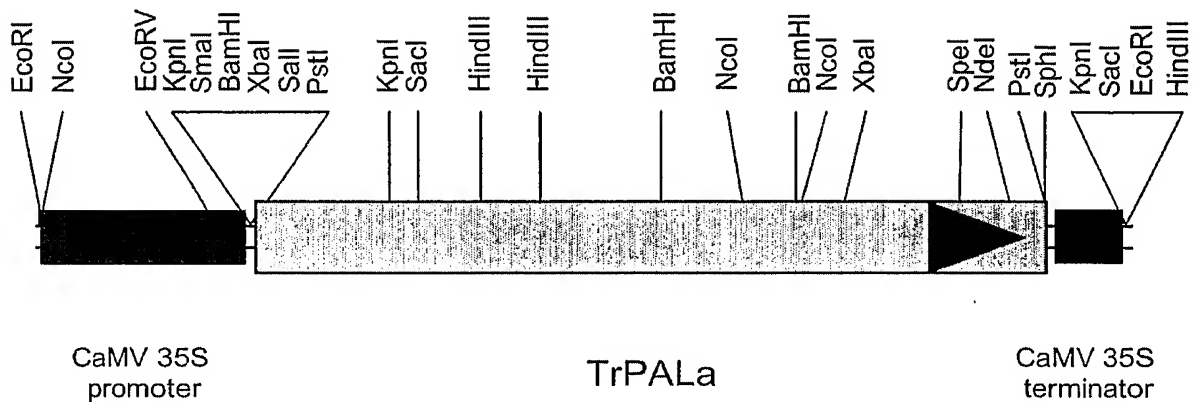
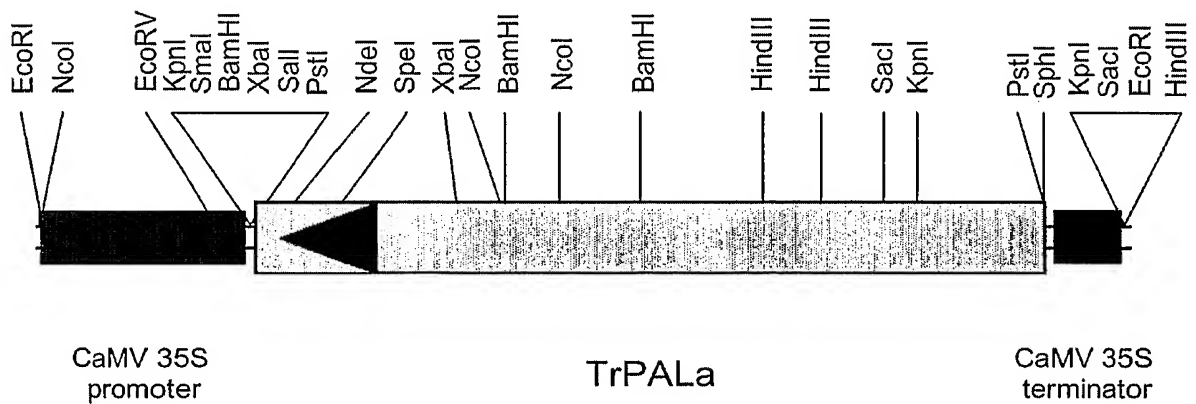
## FIGURE 177

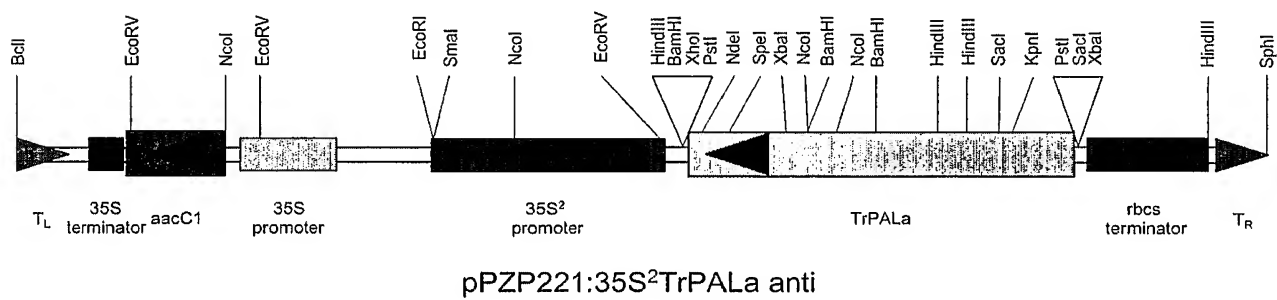
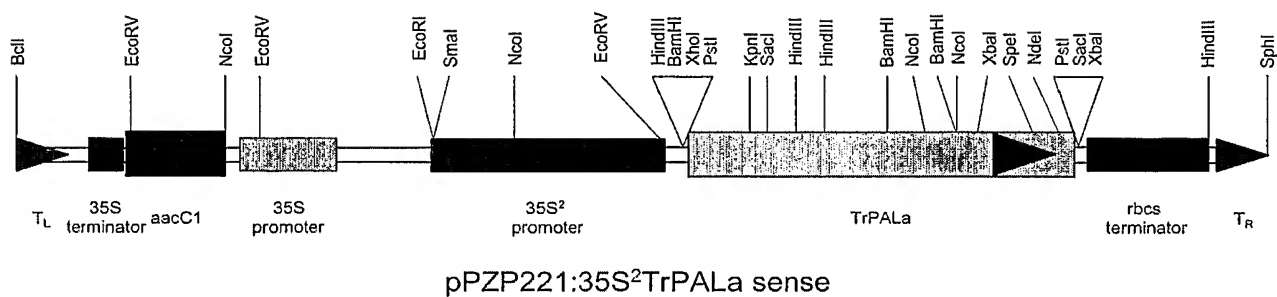


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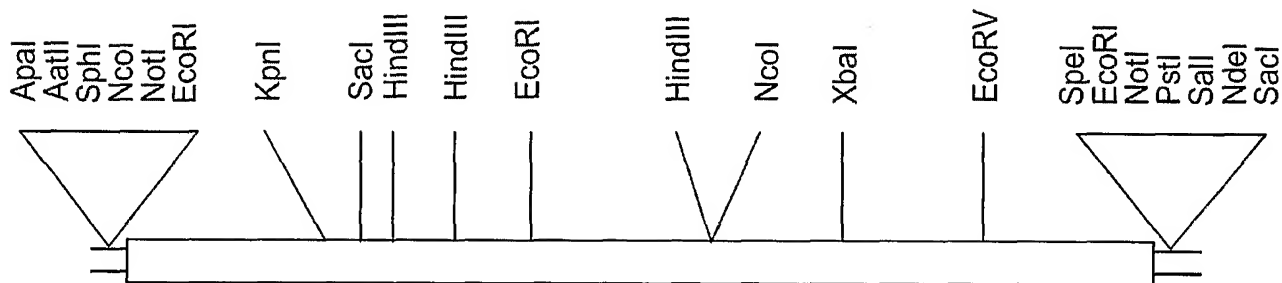
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101 SSDWVMESMN KGTD SYGVPT GFGATSHRRT KQGGALQKEL IRFLNAGIFG  
151 NGTESSHTLP HTATRAAMLV RINTLLQGYS GIRFEILEAI TKLLNNNVTP  
201 CLPLRGTITA SGDLVPLSYI AGLLTGRPNS KAHGPSGEVL NAKQAFQSAG  
251 IDAEFFELQP KEGLALVNGT AVGSGLASIV LFEANILAVL SEVLSAIFAE  
301 VMQ GKPEFTD HLTHKLKHP GQIEAAAIME HILDGSAYVK DAKKLHEMDP  
351 LQKPKQDRYA LRTSPQWLGP LIEVIRFSTK SIEREINSVN DNPLIDVSRN  
401 KALHGGNFQG TPIGVSM DNT RLALASIGKL MFAQFSELVN DFYNNGLPSN  
451 LSASRNPSLD YGFKGSEIAM ASYCSELQYL ANPVTTHVQS AEQHNQDVNS  
501 LGLISSRKTY EAIEILQLMS STFLIALCQA IDLRHLEENL KNSVKNTVSQ  
551 VAKKTLTIGV NGELHPSRFC EKDLLKVVD R EHV FAYIDDP CSATYPLMQK  
601 LRQVLVDHAL VNGESEKNLN TSIFQKIATF EEELKNLCQK RLKVQGLHMK  
651 VEIQQFQTRS MDADLIHSTI L

## FIGURE 178

**252/271****pDH51TrPALa sense****pDH51TrPALa anti****FIGURE 179**

**253/271****FIGURE 180**

**254/271**



**TrPALb**

**FIGURE 181**

# 255/271

```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG AGGAAATAAAA
51  TTCATCATTG TTCCTTATTT CCCACCCAAC ACAACATAAC AAATACATTT
101 CCTCTCCTCT CATCACAAAT ATTACTTTCT ACACCCCCC CTCTCAACTA
151 TTATTAAC TA ACATAATGGA GGAATTACC AATGGCCATG CTGAAGCAAC
201 TTTTGTGCGT ACCAAAAGTG TTGGTGATCC ACTCAACTGG GGTGCAGCCG
251 CGGAGTCGTT GATGGGGAGT CATTTGGATG AGGTGAAGCG TATGGTGGAG
301 GAATACCGTA ATCCATTGGT TAAAATTGGC GGCGAGACGC TTACCATTCG
351 TCAGGTGGCT GGAATTGCTT CTCATGATAG TGGTGTGAGG GTGGAGCTGT
401 CTGAGTCCGC CAGGGCCGGC GTTAAGGCGA GTAGTGGTTG GGTGATGGAC
451 AGCATGAACA ATGGGACTGA TAGTTATGGT GTTACCACTG GTTTCGGCGC
501 CACCTCTCAC CGGAGAACCA AGCAGGGTGG TGCCTTGCAG AAGGAGCTAA
551 TTAGGTTTTT GAATGCCGGA ATATTTGGCA ATGGTACAGA ATCTAATCTG
601 ACCTACCAC ACACAGCAAC CAGAGCTGCA ATGCTTGTGA GAAACAACAC
651 TCTTCTTCAA GGATATTCTG GAATTAGATT TGAAATTTTG GAAAGCTATCA
701 CAAAGCTTCT AAACAACAAC ATTACCCCAT GTTTACCACT TCGTGGTACA
751 ATCACGGCTT CCGGTGATCT CGTCCGCTT TCCTACATTG CCGGTTTGT
801 AACCGGTAGA CCGAACTCCA AAGCCGTTGG ACCCTCCGGA GAAATCTCTCA
851 ATGCAAAAGA AGCTTTTCAA CTGCGCGGCA TTGGTTCTGA GTTTTTTGAA
901 TTGCAGCCAA AAGAAGGTCT TGCTCTTGTT AATGGTACTG CTGTTGGTTC
951 TGGTTTAGCT TCTATTGTTT TGTTTGAAGC AAATGTACTA GCTGTTTTGT
1001 CTGAAGTTAT GTCGGCGATT TTCGCTGAAG TTATGCAAGG GAAACCAGAA
1051 TTCATGATC ATTTGACTCA TAAGTTGAAA CATCACCCCTG GTCAAATTGA
1101 AGCTGCTGCA ATTATGGAAC ATATTTTGGA TGGAAGTGCT TATGTTAAAG
1151 CAGCTAAGAA ATTACACGAA ACCGATCCTT TACAAAAGCC GAAACAAGAT
1201 CGTTATGCAC TTAGAACTTC ACCTCAATGG CTTGGTCCTT TGATTGAAGT
1251 GATAAGATTT TCAACTAAGT CAATTGAGAG AGAAATTAAC TCTGTCAATG
1301 ATAACCCTTT GATTGATGTT TCAAGGAACA AGGCCATTCA CCGTGGTAAT
1351 TTTCAAGGAA CACCTATTGG AGTTTCAATG GATAACACAC GTTTAGCTCT
1401 TGCTTCAATT GGTAAACTCA GTTTTGCTCA ATTCTCTGAA CTGTGTAATG
1451 ATTTTACAA CAACGGGTTA CCTTCGAATC TTAGTGCTAG TAGGAACCCA
1501 AGCTTGGAAT ACGGTTTCAA GGGATCGGAA ATTGCCATGG CTTCTTATTG
1551 TTCTGAGTTA CAATATCTTG CTAATCCTGT CACCACCCAT GTCCAAAGTG
1601 CGGAGCAACA CAATCAAGAT GTTAACCTCT TGGGTTTGAT TTCTTCAAGA
1651 AAAACAAATG AAGCTATTGA GATCCTAAAG CTCATGTCTT CGACATTTCT
1701 GATTGCACTT TGTCAAGCAA TTGATTAAAG GCATTTGGAG GAAATCTGA
1751 GGAACACTGT CAAGAACACG GTAAGCCAAG TAGCGAAGAG AACACTCACC
1801 ACCGGTGTTA ATGGAGAACT TCATCCTTCT AGATTTTGTG AGAAAGATTT
1851 GCTCAAAGTT GTTGATAGGG AGTATGTATT TGCCTATGTC GACGATCCTT
1901 GTCTAGCTAC ATACCCTTTG ATGCAAAAGT TGAGACAAGT GCTTGTGGAT
1951 CATGCATTGG TAAATGCTGA TGGAGAGAAG AATTTGAACA CATCAATCTT
2001 TCAAAAGATT GCAACTTTTG AGGATGAATT GAAAGCTATC TTGCCAAAGG
2051 AAGTTGAAAG TACAAGAACT GCATATGAAA ATGGACAATG TGGAATTTCA
2101 AACAAGATTA AGGAATGCAG GTCTTATCCA TTGTACAAGT TTGTTAGAGA
2151 GGAGTTAGGA ACCGCGTTGC TAACCGGAGA AAAAACGATA TCGCTGGGCG
2201 AAGAGTGTGA CAAATTGTTT ACAGCTATGT GCCAAGGTAA AATTGTTGAT
2251 CCTCTTTTGG AATGCCTTGG AGAGTGGAAT GGTGCTCCTC TACCAATATG
2301 TTAATTAGCA GAATTAATAT GTTTCTTTGA GAAGTGATTT CTTTATATAT
2351 TTGTAGTATA CTATAGTAGT TGCATTGAGA AGCAATTGGT TTGTCTATAA
2401 GCCTATGGAA AATGGCAAAA CAATTTTCTG CTCAAAGCAT CGTTTATTAA
2451 GTTTTCCTTA AAGTGTTAAG GAACTTTAA TTGTTTGTG AATAGAATTT
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2551 TTTGATGTAA ATGGTGTTTT CTCAATTAAT AAATAGTGTT TCAGCCATGA
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2651 TAATCACTAG TGAATTC

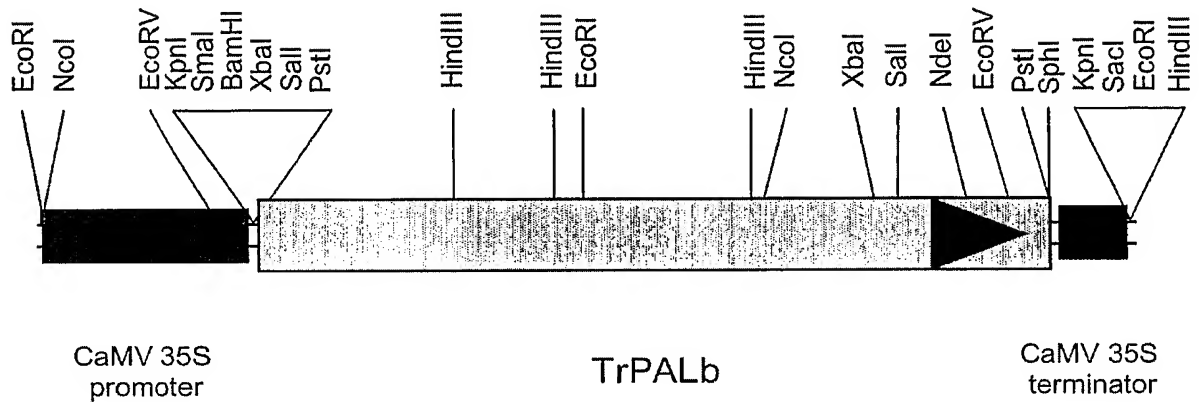
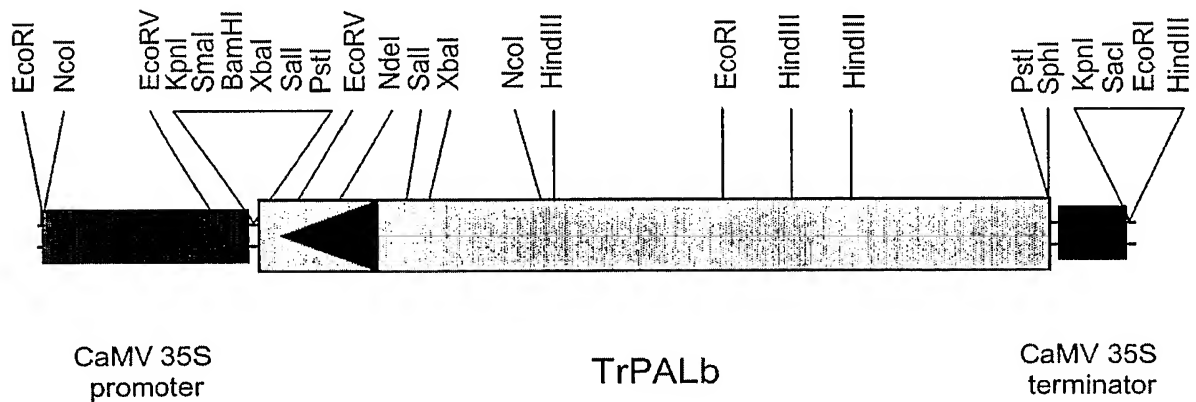
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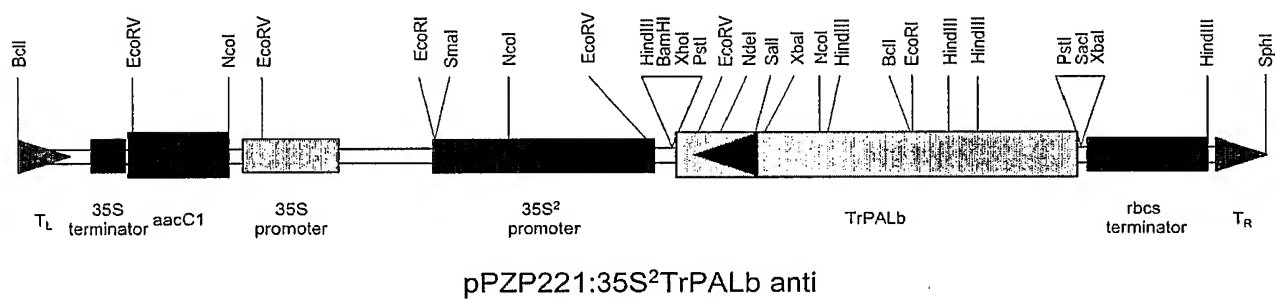
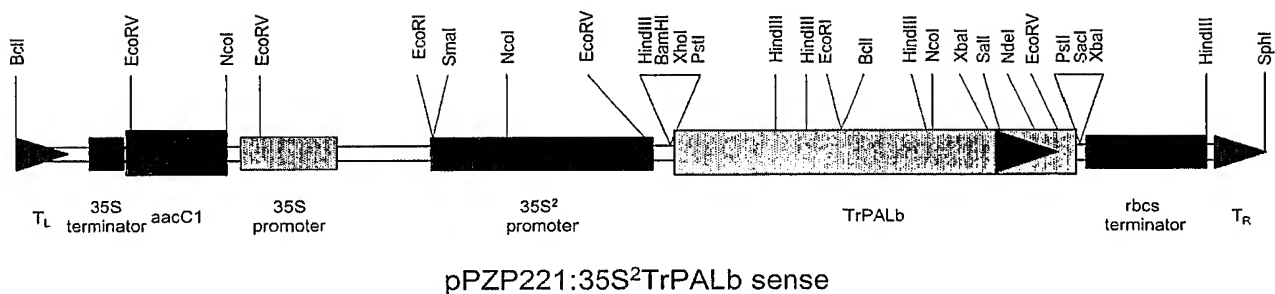
## FIGURE 182

**256/271**

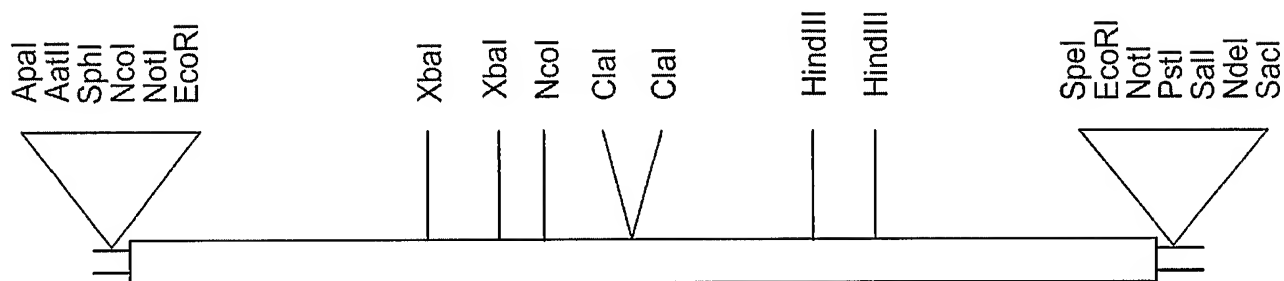
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101 TDSYGVTTGF GATSHRRTKQ GGALQKELIR FLNAGIFGNG TESNCTLPHT  
151 ATRAAMLVRI NTLLQGYSGI RFEILEAITK LLNNNITPCL PLRGTITASG  
201 DLVPLSYIAG LLTGRPNska VGPSGEILNA KEAFQLAGIG SEFFELQPKE  
251 GLALVNGTAV GSGLASIVLF EANVLAVLSE VMSAIFAEVM QGKPEFTDHL  
301 THKLKHHPGQ IEAAAIMEHI LDGSAYVKAA KKLHETDPLQ KPKQDRYALR  
351 TSPQWLGPLI EVIRFSTKSI EREINSVNDN PLIDVSRNKA IHGGNFQGTP  
401 IGVSM DNTRL ALASIGKLMF AQFSELVNDF YNNGLP SNLT ASRNPSLDYG  
451 FKGSEIAMAS YCSELQYLAN PVTTHVQSAE QHNQDVNSLG LISSRKTNEA  
501 IEILKLMSST FLIALCQAID LRHLEENLRN TVKNTVSQVA KRTLTTGVNG  
551 ELHPSRFCEK DLLKVVDREY VFAYVDDPCL ATYPLMQKLR QVLVDHALVN  
601 ADGEKNLNTS IFQKIATFED ELKAILPKEV ESTRTAYENG QCGISNKIKE  
651 CRSYPLYKFV REELGTALLT GEKTISLGEE CDKLFTAMCQ GKIVDPLEEC  
701 LGEWNGAPLP IC

**FIGURE 183**

**257/271****pDH51TrPALb sense****pDH51TrPALb anti****FIGURE 184**

**258/271****FIGURE 185**



**259/271****TrPALf****FIGURE 186**

# 260/271

```

1  GAATTCACTA GTGATTAAGC AGTGGTAACA ACGCAGAGTA CGCGGGGAGG
51 AAATAAATTC ATCATTGTTC ATTATTTCCC ACCCAACACA ACATAACAAA
101 TACATTATTC TCTCCTCTGA TCACAATTAT TACTTTCTAC ACCCTCCTCT
151 CAACTATTAT TAACTAGCAT AATGGAGGGA ATTACCAATG GCCATGCTGA
201 AACAACTTTT TGCGTGACCA AAAGTGTGGG TGATCCACTC AACTGGGGTG
251 CAGCCGCGGA GTCGTTGACG GGGAGTCATT TGGATGAGGT GAAGCGTATG
301 GTGGAGGAGT ACCGTAATCC GTTGGCTAAA ATTGGCGGCG AGACGCTTAC
351 CATTGCTCAG GTGGCTGGAA TTGCTTCTCA TGATAGTGGT GTGAGGGTGG
401 AGCTGTCCGA GTCCGCAAGG GCCGGCGTTA AGGCGAGTAG TGATTGGGTG
451 ATGGATAGCA TGAACAATGG GACTGATAGT TACGGTGTTA CCACCGGTTT
501 TGGTGCCACC TCTCACC CGGA GAACCAAGCA GGGTGGTGC TGCAGAAGG
551 AGCTAATTAG GTTTTTGAAT GCTGGAATAT TTGGCAATGG TACAGAATCT
601 AACTGTACAC TACCACACAC AGCAACTAGA GCTGCAATGC TTGTGAGAAT
651 CAACACTCTT CTTCAAGGGT ACTCTGGTAT TAGATTTGAA ATTTTGGGAG
701 CTATCACAAA GCTTCCAAAC AACAACATTA CCCCATGTTT ACCACTTCGT
751 GGTACAATCA CGGCTTCCGG TGATCTTGTT CCGCTTTCCT ACATTGCCCG
801 TTTGTTAACC GGAAGACCCA ACTCCAAAGC AGTTGGACCT TCCGGAGAAA
851 TTTTGAGTGC TAAAGAAGCT TTCAACTCG CCGGCATTGG TTCTGAGTTT
901 TTTGAATTGC AACCAAAAGA AGGTCTTGCT CTTGTTAATG TACTGTGCTG
951 TGGCTCTGGT TTAGCTTCTA TTGTTCTGTT TGAAGCAAAT GTACTAGCTG
1001 TTTTATCCGA AGTTATGTCG GCGATTTTCG CTGAAGTTAT GCAAGGGA
1051 CCGGAATTTA CCGATCATTT GACTCATAAG TTGAAACATC ACCCTGGTCA
1101 AATTGAAGCT GCTGCAATTA TGGAACATAT TTTGGATGGA AGTGCTTATG
1151 TTAAAGCAGC TAAGAAGTTA CACGAAACCG ATCCTTTACA AAAACCGAAA
1201 CAAGATCGTT ATGCACTTAG CCAATTCGAT TGAAAGAGAA ATTAATCGG
1251 TGAAGTGATA AGATTTTCAA CCATTTGATC GATGTTTCAA GGAACAAGGC CATTCATGGT
1301 TCAACGACAA CCCTTTGATC GATGTTTCAA GGAACAAGGC CATTCATGGT
1351 GGTAAC TTTC AAGGAACACC TATTGGAGTT TCAATGGATA ACACACGTTT
1401 AGCTCTTGCT TCAATTGGTA AACTCATGTT TGCTCAATTC TCTGAACTTG
1451 TTAATGATTT TTACAACAAC GGGTGTGCCTT CGTATCTTAC TGCTAGTAGG
1501 AACCCGAGCT TGGACTATGG TTTCAAGGGA TCGGAAATTG CCATGGCTTC
1551 GTATTGTTCC GAGTTACAAT ATCTTGCTAA TCCTGTCACC ACCCATGTCC
1601 AAAGTGCCGA GCAACACAAC CCAATGTTA ACTCTTTGGG TTTGATTTCCT
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1701 TTTCTTGATT GCATTATGTC AAGCAATCGA CTTAAGGCAC TTGGAGGAAA
1751 ATCTCAGGAA CACCGTCAAG AACACGGTAA GCCAAGTAGC GAAGAGAACA
1801 CTCACCACCG GCGTCAACGG AGAACTTCAT TCTTCTAGAT TTTGTGAGAA
1851 AGATTTGCTT AAAGTTGTTG ATAGGGAGTA TGTATTTGCC TATGCCGACG
1901 ATCCTTGCTC AGCTACATAC CCTTTGATGC AAAAGTTGAG ACAAGTGCTT
1951 GTGGATCATG CATTGGTAAA TGTTGATGGA GAGAAGAATT TGAACACATC
2001 AATCTTTCAA AAGATTGCAA CTTTTGAGGA TGAGTTGAAA GCTATTTTGC
2051 CAAAGGAGGT TGAAAGTACA AGAACTGCAT ATGAAAATGG ACAATGTGGA
2101 ATTTCAAACA AGATTAAGGA ATGCAGGTCT TATCCATTGT ACAAGTTTGT
2151 TAGAGAGGAG TTAGGAACCG CGTTGCTAAC CGGAGAAAAA ACTATATCGC
2201 CGGGCGAAGA GTGCGATAAA TTGTTTACAG CTATGTGCCA AGGTAAAATT
2251 GTTGATCCTC TTATGGAATG CCTCGGAGAG TGGAATGGTG CTCCTCTACC
2301 AATATGTTAA TTAGCATAAT ATGTTTCTT TGAGAAGTGA TTACTTTATA
2351 TATTTGTAGT ATACTATAGT AGTTGCATTG AGAAGAAATT GGTTTGTTTA
2401 TAAGCCTATG GAAAATGGCA AATCAATTTT CTGCTCAAAG CATCGTTTAT
2451 TAAGTTTTCC TTAAAGTGTT AAGGAACTTT TAATTGTTTT TGTAATAGAA
2501 TTTCATTTGT TTGCCACAAC TTTGGGTGCA AATATCACAT GATACATGTG
2551 GTGTTTGATG TAAATGGTGT TTTTTCAATA AATAAATAGT GTTTCAACTA
2601 AAAAAAAAAA AAAAAAAAAA AAAAAAAAAA TACTCTGCGT TGTTACCACT
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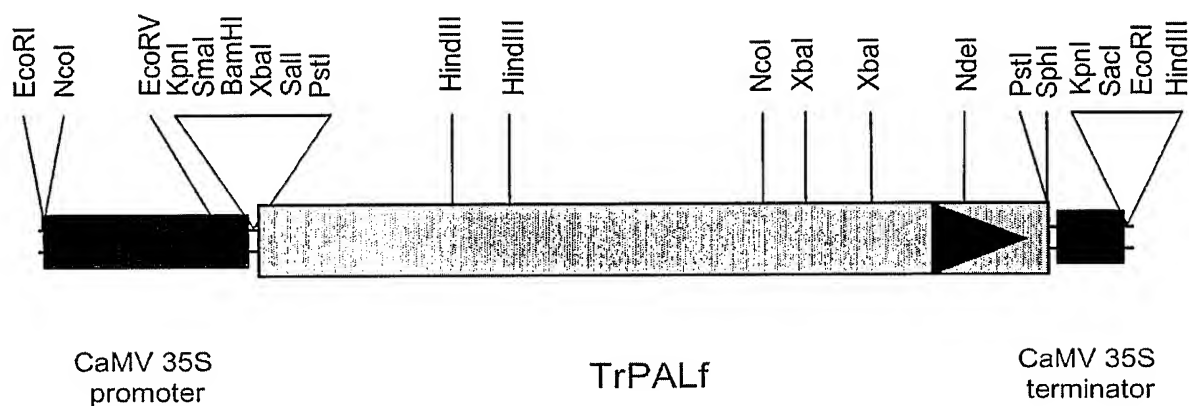
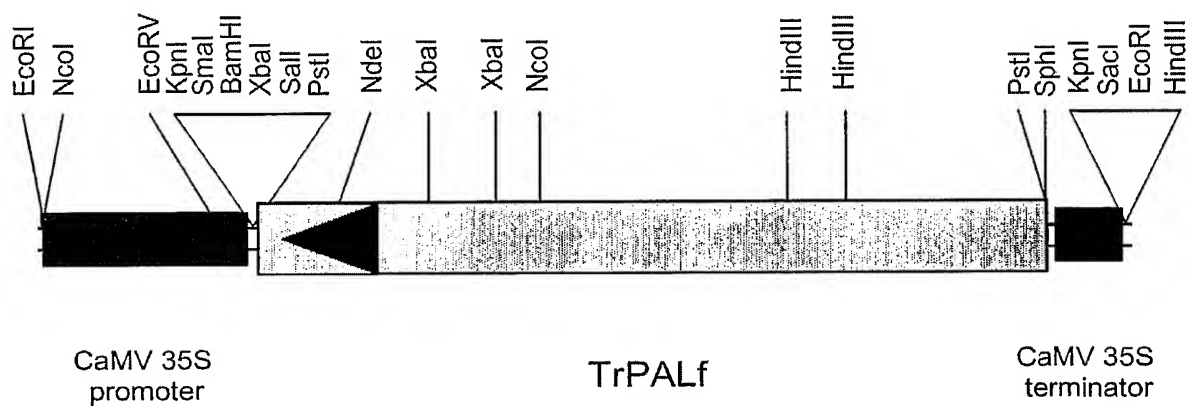
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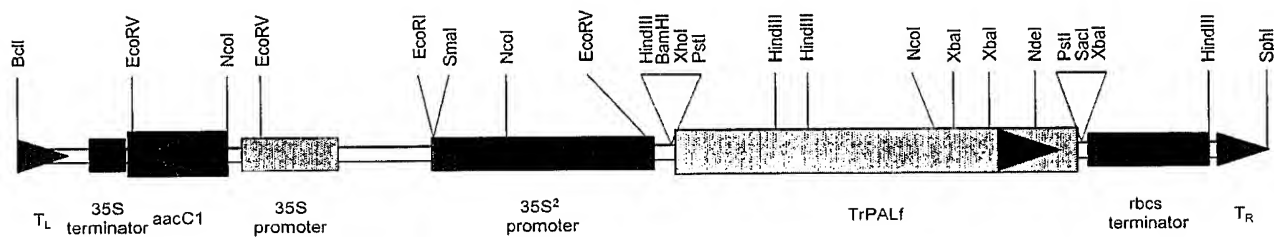
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# 261/271

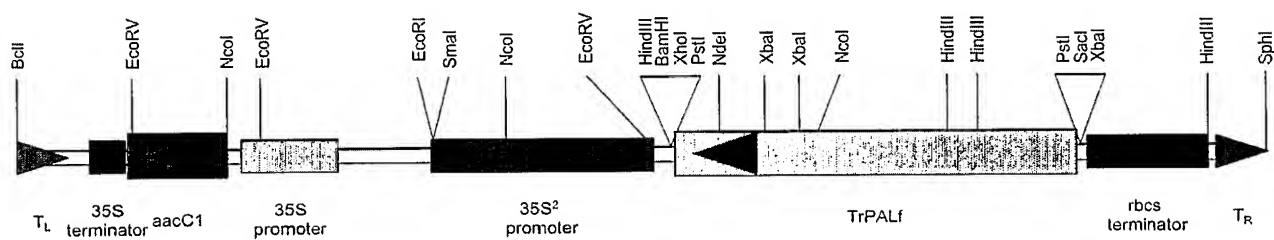
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101 TDSYGVTTGF GATSHRRTKQ GGALQKELIR FLNAGIFGNG TESNCTLPHT  
151 ATRAAMLVRI NTLLQGYSGI RFEILEAITK LPNNNITPCL PLRGTITASG  
201 DLVPLSYIAG LLTGRPNska VGPSGEILSA KEAFQLAGIG SEFFELQPKE  
251 GLALVNGTAV GSGLASIVLF EANVLAVLSE VMSAIFAEVM QGKPEFTDHL  
301 THKLKHHPGQ IEAAAIMEHI LDGSAYVCAA KKLHETDPLQ KPKQDRYALR  
351 TSPQWLGPLI EVIRFSTKSI EREINSVNDN PLIDVSRNKA IHGGNFQGTP  
401 IGVSM DNTRL ALASIGKLMF AQFSELV NDF YNNGLP SYLT ASRNPSLDYG  
451 FKGSEIAMAS YCSELQYLAN PVTTHVQSAE QHNQDVNSLG LISSRKTNEA  
501 IEILKLMSST FLIALCQAID LRHLEENLRN TVKNTVSQVA KRTLTTGVNG  
551 ELHSSRFCEK DLLKVVDREY VFAYADDPCL ATYPLMQKLR QVLVDHALVN  
601 VDGEKNLNTS IFQKIATFED ELKAILPKEV ESTRTAYENG QCGISNKIKE  
651 CRSYPLYK FV REELGTALLT GEKTISP GEE CDKLFTAMCQ GKIVDPLMEC  
701 LGEWNGAPLP IC

## FIGURE 188

**262/271****pDH51TrPALf sense****pDH51TrPALf anti****FIGURE 189**

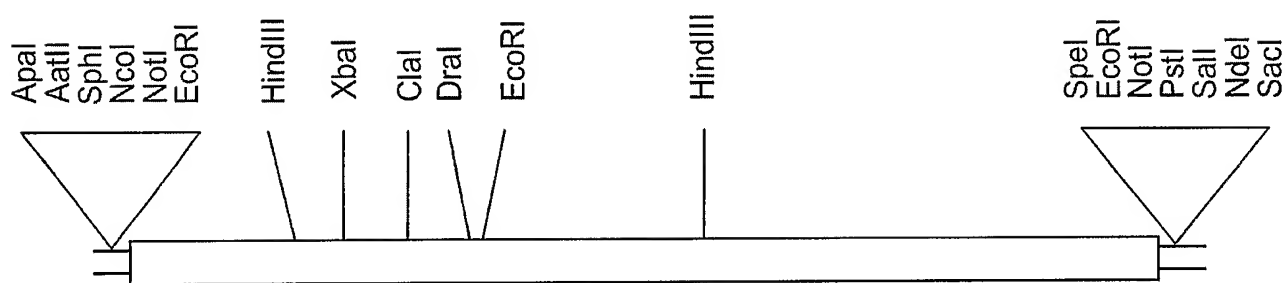
**263/271**

pPZP221:35S²TrPALf sense



pPZP221:35S²TrPALf anti

**FIGURE 190**

**264/271****TrVRa****FIGURE 191**

# 265/271

```

1  GAATTCGATT AAGCAGTGGT AACAAACGCAG AGTACGCGGG ATAGTAGTAG
51 TTGAGAAAAA ATACACAAAT AAAGTAAACA CTATCATAGA AAGAGAGTCA
101 AAAATGGCTG AAGGAAAAGG AAGGGTTTGT GTTACTGGAG GAACAGGTTT
151 TCTTGGTTCA TGGATCATCA AGAGTCTTCT TGAAAAATGGA TACTCTGTTA
201 ATACCACTAT TAGAGCTGAT CCAGAACGTA AGAGGGATGT AAGCTTCCTA
251 ACAAATCTAC CCGGCGCATC CGAAAGGCTA CATTTCTTCA ACGCCGATCT
301 AGACGACCCA GAGAGTTTCA ACGAAGCAAT TGAAGGTTGT GTCGGGATAT
351 TCCACACCGC TTCACCAATC GATTTCGCCG TGAGTGAGCC AGAAGAAATA
401 GTGACAAAAA GAACAGTGGA TGGAGCATTG GGAATTTTAA AAGCATGTGT
451 GAATTCAAAG ACAGTGAAGA GATTTATTTA CACTTCAAGT GGTTCCTGCTG
501 TTTCATTTCAA TGGAAAAAAC AAAGATGTTT TGGATGAGAG TGATTGGAGT
551 GATGTTGATT TGCTTAGAAG TGTTAAACCA TTTGGTTGGA GTTATGGTGT
601 TTCAAAGACT TTGGCTGAGA AAGCAGTGCT TGAATTTGGT CAACAAAATG
651 GGATTGATGT TGTTACTTTG ATTCTTCCTT TTATTGTTGG AAGTTTTGTT
701 TGTCTAAGC TTCCTGATTC TGTTGAGAAA GCTCTTGTTT TGGTACTAGG
751 CAAAAAGGAA CAAATTGGTA TTATAAGTTT CCACATGGTA CATGTGGATG
801 ATGTGGCTAG AGCACATATT TATCTACTTG AGAATCCTGT TCCAGGAGGT
851 AGATATAATT GTTCACCATT CTTTGTATCT ATTGAAGAA TGTCACAAC
901 TCTTTCAGCC AAATATCCAG AATATCAAA ACTATCAGTA GATGAGTTGA
951 AGGAAATCAA AGGTGCAAGG TTGCCAGATT TGAACTCGAA AAAGCTCGTG
1001 GACGCTGGTT TTGAGTTTAA GTATAGTGTC GGTGATATGT TCGATGATGC
1051 GATTCAATGC TGCAAGGAAA AAGGCTATCT CTAAGTATGT GTTTGAAAAA
1101 AATTCCATGA AGCTGAGAAA ACAATAATAT GCCTAAAATC AATGATGGCT
1151 AATGAAATGT ACAAGTTTAT GCATAAAGTT ATTTGTGATG AATCAAATAA
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1251 AAAAAGTACT CTGCGTTGTT ACCACTGCTT AATCACTAGT GAATTC

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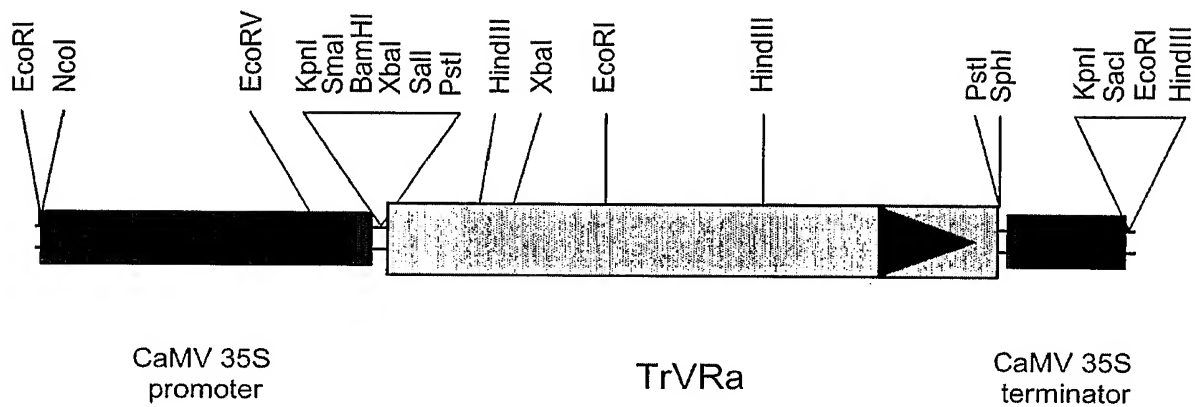
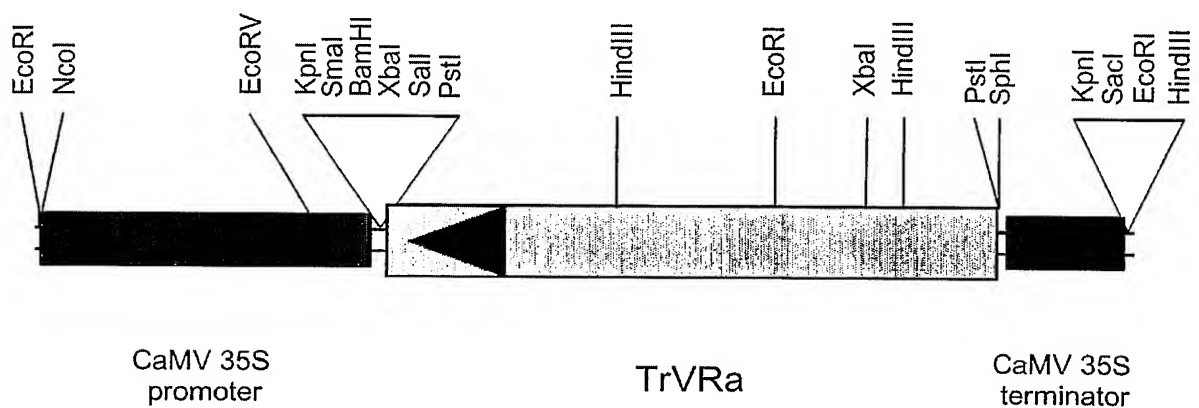
## FIGURE 192

# 266/271

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1  MAEGKGRVCV TGGTGFLGSW IIKSLENGY SVNTTIRADP ERKRDVSFLT
51  NLPGASERLH FFNADLDDPE SFNEAIEGCV GIFHTASPID FAVSEPEEIV
101 TKRTVDGALG ILKACVNSKT VKRFIYTSSG SAVSFNGKNK DVLDESDWSD
151 VDLLRSVKPF GWSYGVSKTL AEKAVLEFGQ QNGIDVVTLI LPFIVGSFVC
201 PKLPDSVEKA LVLVLGKKEQ IGIISFHMVH VDDVARAHY LLENPVPGGR
251 YNCSPFFVSI EEMSQLLSAK YPEYQILSVD ELKEIKGARL PDLNSKKLVD
301 AGFEFKYSVG DMFDDAIQCC KEGYL
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## FIGURE 193



**267/271****pDH51TrVRa sense****pDH51TrVRa anti****FIGURE 194**

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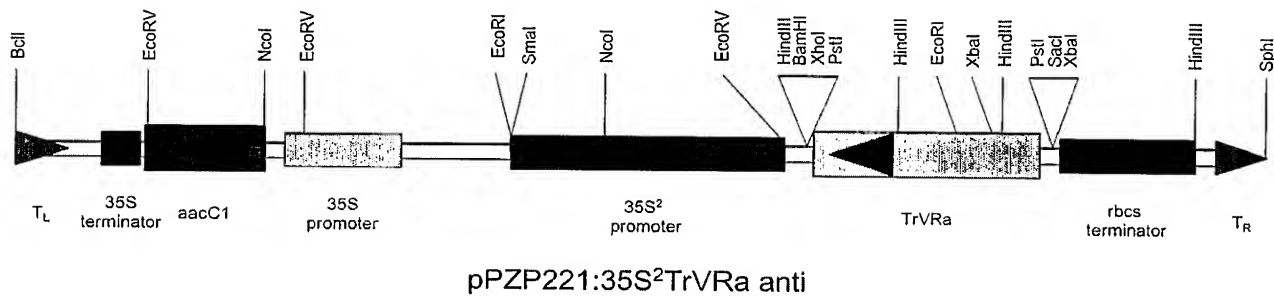
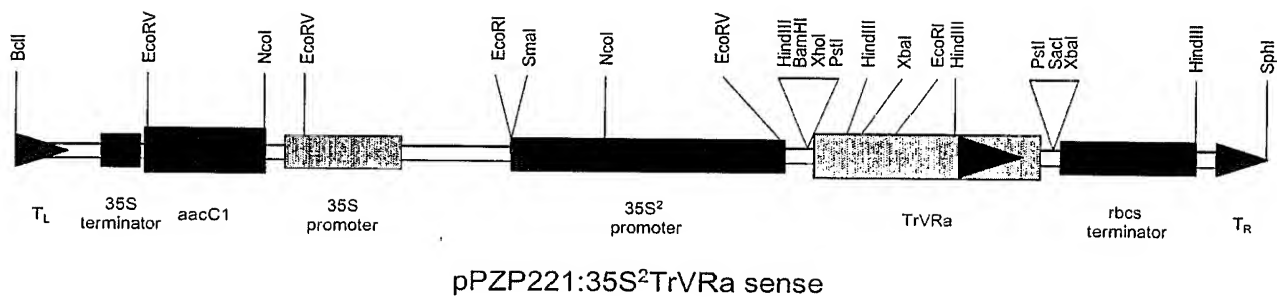
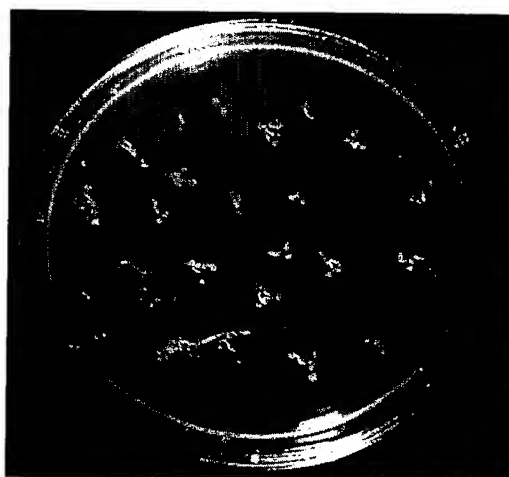
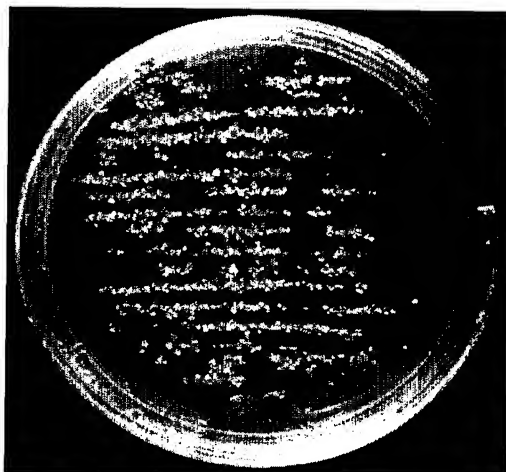
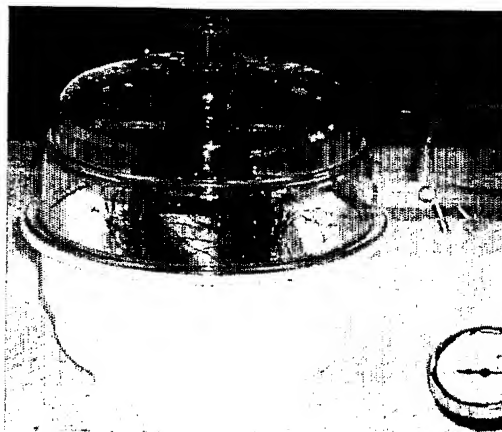
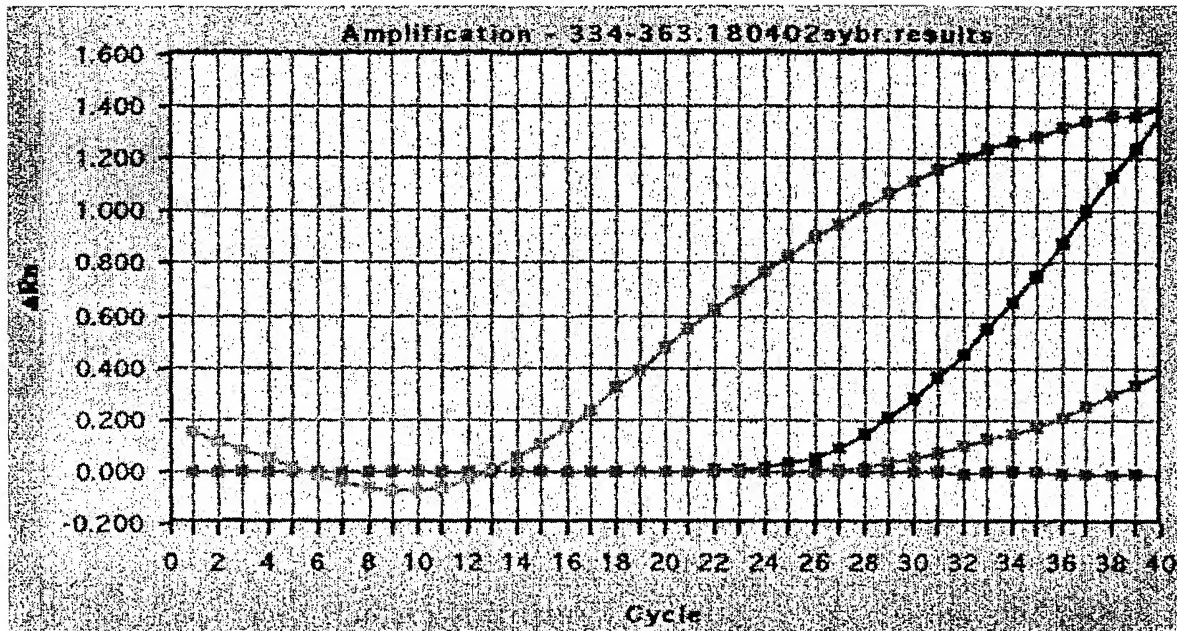
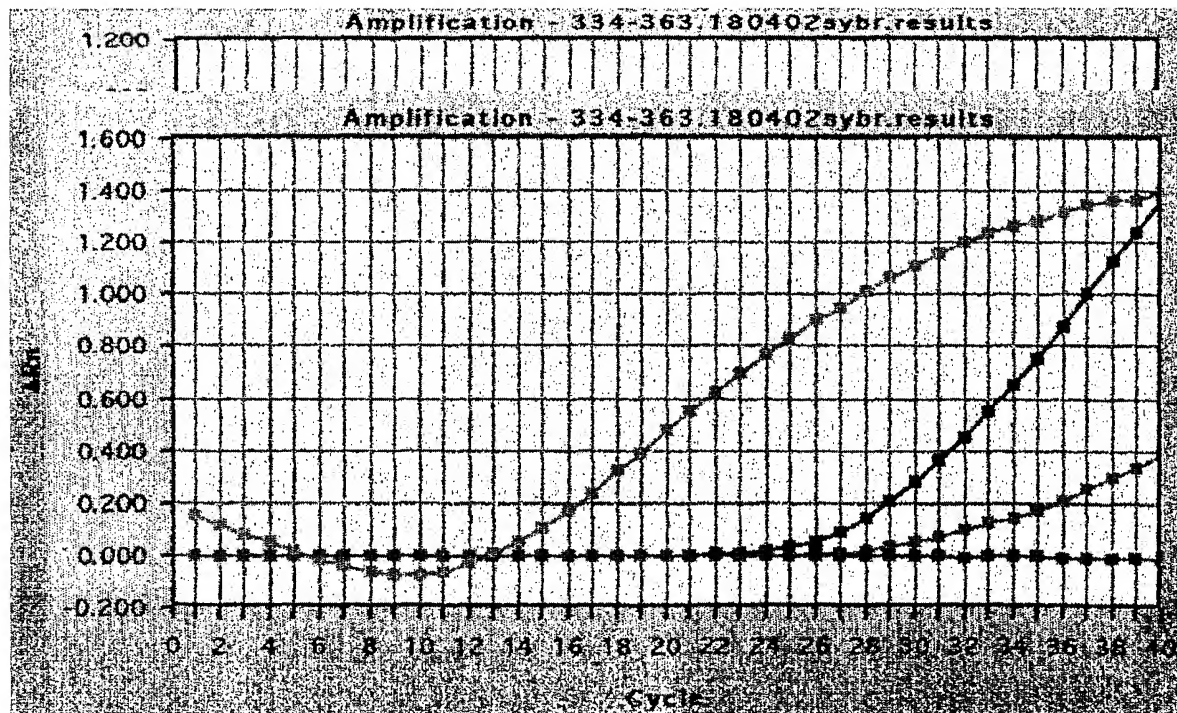


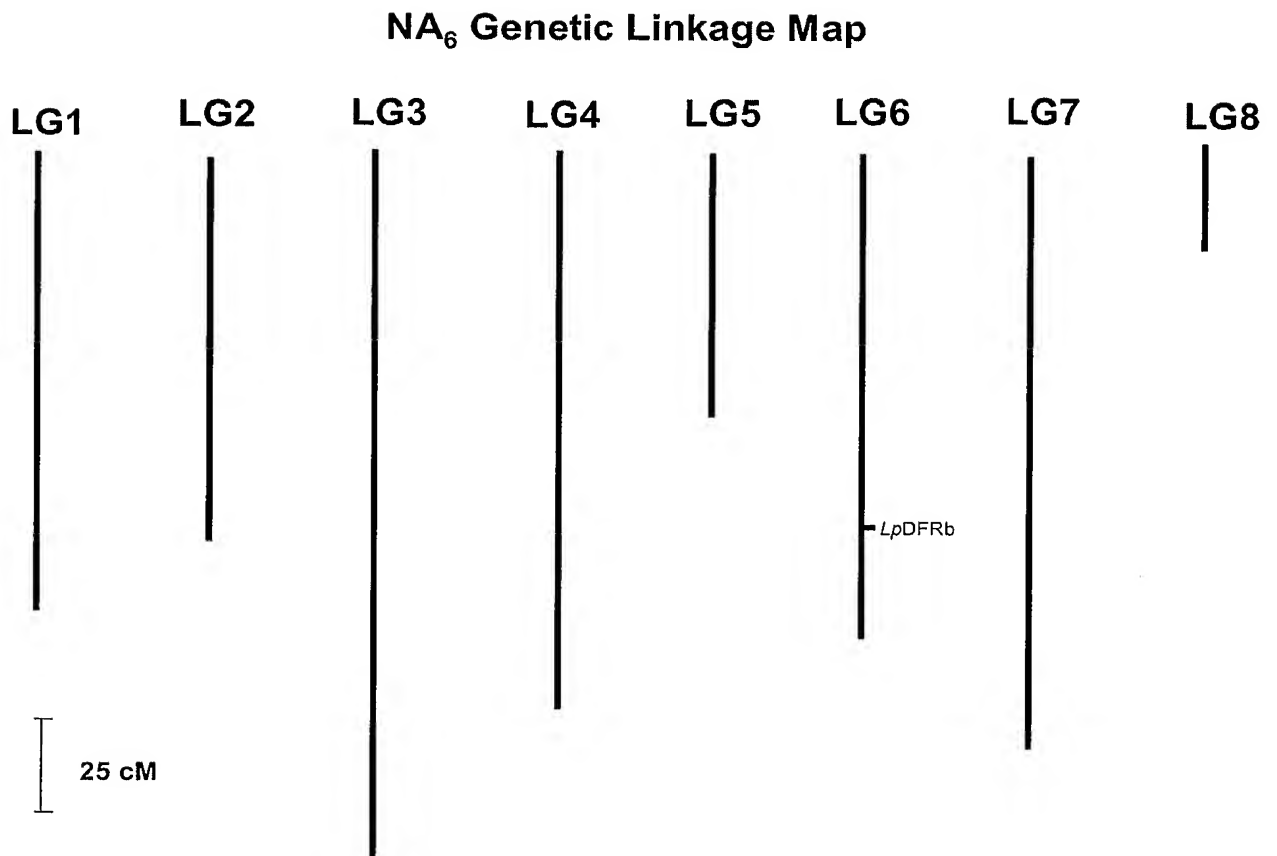
FIGURE 195

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**FIGURE 196**

**270/271****D****E****FIGURE 196 (cont)**

**271/271****FIGURE 197**

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AgResearch Limited

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Gly Glu Arg Gly Leu Thr Ile Glu Gly Asn Phe Ile Lys Phe Thr Ala  
 35 40 45

Ile Gly Val Tyr Leu Glu Asp Val Ala Val Ala Ser Leu Ala Thr Lys  
 50 55 60

Trp Lys Gly Lys Ser Ser Glu Glu Leu Leu Glu Thr Leu Asp Phe Tyr  
 65 70 75 80

Arg Asp Ile Ile Ser Gly Pro Phe Glu Lys Leu Ile Arg Gly Ser Lys  
 85 90 95

Ile Arg Glu Leu Ser Gly Pro Glu Tyr Ser Arg Lys Val Asn Glu Asn  
 100 105 110

Cys Val Ala His Leu Lys Ser Val Gly Thr Tyr Gly Asp Ala Glu Val  
 115 120 125

Glu Ala Met Gln Lys Phe Val Glu Ala Phe Lys Pro Ile Asn Phe Pro  
 130 135 140

Pro Gly Ala Ser Val Phe Tyr Arg Gln Ser Pro Asp Gly Ile Leu Gly  
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120

ccggctgtgg ttacttctcc ggccaccggt aagtcattt ttcttggtgg tgcaggggag  
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120

acttctccgg tcaacggtaa gtcattttt cttggtggtg caggggagag aggtttgact  
180

attgaaggaa acttcatcaa gttcactgcc ataggagtat atttgaaga ttagcagtg  
240

gcttcacttg ccactaaatg gaagggcaaa tcctctgaag agttgcttga gacccttgac  
300

ttctacagag atatcatttc aggaccattt gagaagttga ttcgaggatc gaagattagg  
360

gaattgagtg gtcctgagta ctcaaggaag gttaatgaaa actgtgtggc acacttaaaa  
420

tctgttgga cttacggaga tgcagaagtt gaagctatgc aaaaatttgt tgaagccttc  
480

aagcctatta attttccacc tggtgcctct gttttttaca ggcaatcacc tgatggaata  
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catattttct tgggtggtgca ggggagagag gtttgactat tgaaggaaac ttcacaaagt  
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tcactgccat aggagtatat ttggaagatg tagcagtggc ttcacttgcc actaaatgga  
240  
agggtaaatc ctctgaggag ttgcttgaga ctcttgactt ctatagagac atcatttcag  
300  
gaccctttga aaagttgatt cgaggatcga agattagga attgagtggc cctgagtact  
360  
caaggaaggt taatgaaaac tgcgtggccc acttaaaatc tgttgggact tatggagatg  
420  
ctgaagctga agctatgcaa aaatttggtg aagccttcaa gcctattaat tttccacctg  
480  
gtgcctctgt tttttacagg caatcacctg atggaatatt agggccttagt ttctctcaag  
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tcatattttc ttggtggtgc aggggagaga ggtttgacta ttgaaggaaa cttcatcaag  
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gtgattactt ctccggtcaa cggtaagtca tattttcttg gtggtgcagg ggagagaggt  
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ttgactattg aaggaaactt catcaagttc actgccatag gagtatattt ggaagatgta  
240  
gcagtggctt cacttgccac taaatggaag ggcaaactct ctgaagagtt gcttcagacc  
300  
cttgacttct acagagatat catttcagga ccatttgaga agttgattcg aggatccaag  
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480  
gccttcaagc ctattaattt tccacctggt gcctctgttt tttacaggca atcacctgat  
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120

gtcgagaacc ttgaattccc ggcggtgatt acttctccgg tcaacggtaa gtcataatatt  
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cttggtggtg caggggagag aggttngact attgaaggaa acttcatcaa gttcactgcc  
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Val Ile Thr Ser Pro Val Asn Gly Lys Ser Tyr Phe Leu Gly Gly Ala  
20 25 30

Gly Glu Arg Gly Xaa Thr Ile Glu Gly Asn Phe Ile Lys Phe Thr Ala  
35 40 45

Ile Gly Val Tyr Leu Glu Asp Val Ala Gly Ala Ser Leu Ala Thr Lys  
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120

gtcgagaacc ttgaattccc ggcggtgatt atttctccgg tcaacggtaa gtcatatattt  
180

cttggtggtg caggggagag aggtttgact attgaaggaa acttcatcaa gttcactgcc  
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ataggagtat atttgaaga thtagcaggg gttcacttg ccactaaatg gaagggnaga  
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120  
tcgagaacct tgaattcccg gcggtgatta cttctccggt caacggtaag tcatattttc  
180  
ttggtggtgc aggggagaga ggttngacta ttgaaggaaa cttcatcaag ttcactgcca  
240  
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120

tggtgggact tatggagatg cagaagctga agctatgcaa aaatttggtg aagccttcaa  
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gcctattaat ttccacctg gtgcctctgt tttttacagg caatcacctg atggaatatt  
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274

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Gly Pro Glu Tyr Ser Arg Lys Val Asn Glu Asn Cys Val Ala His Leu  
20 25 30

Lys Ser Val Gly Thr Tyr Gly Asp Ala Glu Ala Glu Ala Met Gln Lys  
35 40 45

Phe Val Glu Ala Phe Lys Pro Ile Asn Phe Pro Pro Gly Ala Ser Val  
50 55 60

Phe Tyr Arg Gln Ser Pro Asp Gly Ile Leu Gly Val Ser Ile Ala Asn  
65 70 75 80

Ser Phe Phe Leu Thr Ile Leu Ile Arg Val Arg Phe Asp Cys  
85 90

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120

tggtgggact tatggagatg cagaagctga agctatgcaa aaatttggtg aagccttcaa  
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gcctattaat tttccacctg gtgcctctgt tttttacagg caatcacctg atggaatatt

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aggggtagt attgccaatt cattttttttt aact  
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120

tatggagatg cagaagctga agctatgcaa aaatttggtg aagccttcaa gcctattaat  
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tttccacctg gtgcctctgt tttttacagg caatcacctg atggaatatt aggggtagt  
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120

aatctattcc ctctaccgt cacaccaccg ggatccacca acaatttctt cctcggcgggt  
180

gcaggagagc ggggtcttca aattcaagac aaatttgtca aattcaccgc tattggtggt  
240

tatctacagg acattgctgt tccttacctc gccactaaat ggaagggtaa gactgctcaa  
300

gagctaacgg aaactgttcc tttcttcagg gacatcgta caggctcatt tgagaaattt  
360

atgcaggtga caatgatctt gccattgact gggcaacaat actcagagaa agtgtcagaa  
420

aattgtgtag ctatttgga gtctcttggg atttataccg acgaagaagc caaagcaatt  
480

gagaagnntg tttctgtctt caaagangaa acattcccac caggctcctc tctccttttc  
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acagnattac ccaaaggatt aggatcacta acgataagnt tctctaaaga tggatccatt  
600

ccagagaccg agtctgcagt tatagagaat aagctactct cacaagctgt gctngagtcg  
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atgatagggg cacacgggtgt ctcccttgca gcaaaacaga gttttggcca ccaggntanc  
720

cgagntattc aacgaggntg gctgatgcct agcaacttga tnatatcaac aaaacgaaaa  
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Gly Ala Gly Glu Arg Gly Leu Gln Ile Gln Asp Lys Phe Val Lys Phe  
 35 40 45

Thr Ala Ile Gly Val Tyr Leu Gln Asp Ile Ala Val Pro Tyr Leu Ala  
 50 55 60

Thr Lys Trp Lys Gly Lys Thr Ala Gln Glu Leu Thr Glu Thr Val Pro  
 65 70 75 80

Phe Phe Arg Asp Ile Val Thr Gly Pro Phe Glu Lys Phe Met Gln Val  
 85 90 95

Thr Met Ile Leu Pro Leu Thr Gly Gln Gln Tyr Ser Glu Lys Val Ser  
 100 105 110

Glu Asn Cys Val Ala Ile Trp Lys Ser Leu Gly Ile Tyr Thr Asp Glu  
 115 120 125

Glu Ala Lys Ala Ile Glu Lys Val Ser Val Phe Lys Glu Thr Phe Pro  
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120  
aaagggcaga aggccctgca acaatcttgg ccattggcac tgcaaatacca gcaaaccgtg  
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ttgatcagag cacatatacct gatttctact tcaaaatcac taacagtga cataaagtgt  
240  
agctcaaaga gaaattccag cgcattgtgtg ataaatctat gatcaagagc agatacatgt  
300  
atctaacaga agagattttg aaagaaaatc ctagtctttg tgagcacatg gcaccttcac



360

tggatgctag gcaagacatg gtggtggttg aggtacctag acttgggaag gaggctgcag  
420

tgaaagctat aaaagaatgg ggtcaaccaa agtcaaagat tactcactta atcttttgca  
480

ccacaagtgg tgttgacatg cctgggtgctg attaccaact cacaaaactc ttaggtcttc  
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120

caaagggcag aaggccctgc aaccatttta gccattggta ctgcaaatcc agcaaaccgt  
180

gtagaccaga gcacatatcc tgattttctac ttcaaaatca ctaacagtga gcataagggt  
240

gagcttaaag agaaattcca gcgcatgtgt gataaatcta tgatcaagag cagatacatg  
300

tatctaacag aagagatttt gaaagaaaat cctagtcttt gtgaatacat ggcaccttca  
360

ttggatgcta ggcaagacat ggtgggtggt gaggtacctt gactaggaaa ggaggctgca  
420

gtcaaggcca ttaaagaatg ggtcaacca aagtcaaaga ttactcactt aatcttttgc  
480

accacaagtg gtgtagacat gcctgggtgct gattaccaac tcacaaaact cttgggactt  
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120

gccctgcaac cattttggcc attggcactg caaatccacc aaaccgtggt gagcagagca  
180

catatcctga tttctacttc aaaattacaa acagtgagca caagactgag ctcaaagaga  
240

agttccaacg catgtgtgac aaatccatga tcaagagcag atacatgtat ctaacagaag  
300

agattttgaa agaaaatcct agtctttgtg aatacatggc accttcattg gatgctaggc  
360

aagacatggt ggtgggtgag gtacctagac ttgggaagga ggctgcagtc aaggccatta  
420

aagaatgggg tcaaccaaag tcaaagatta ctacttaat cttttgcacc acaagtgggtg  
480

ttgacatgcc tgggtgctgat taccaactca caaaactctt aggtcttcgc ccatatgtga  
540

aaaggtatat gatgtaccaa caagggtgtt ttgcaggagg cacgggtgctt cgtttggcaa  
600

aagatttggc cgagaacaac aaagggtgctc gtgtgctagt tgtttgttct gaagtcaccg  
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120

ctgaaggccc tgcaaccatt ttggccattg gactgcaaa tccaccaaac cgtgttgagc  
180

agagcacata tcctgatttc tacttcaaaa ttacaaacag tgagcacaag actgagctca  
240

aagagaagtt ccaacgcatg tgtgacaaat ccatgatcaa gagcagatac atgtatctaa  
300

cagaagagat tttgaaagaa aatcctagnc tttgtgaata catgncacct tcattggatg  
360

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gccctgcaac cattttggcc attggcactg caaatccacc aaaccgtggt gagcagagca  
180  
catatcctga tttctacttc aaaattacaa acagtgcagca caagactgag ctcaaagaga  
240  
agttccaacg catgtgtgac aaatccatga tcaagagcag atacatgtat ctaacagaag  
300  
agattttgaa agaaaatcct agtctttgtg aatacatggc accttcattg gatgctaggc  
360  
aagacatggt ggtgggtgag gtacctanac ttgggaagga ggctgcannc aaggccatta  
420  
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120

ctgaaggccc tgcaaccatt ttggccattg gcaactgcaaa tccaccaaac cgtgttgagc  
180

agagcacata tcctgatttc tacttcaaaa ttacaaacag tgagcacaag actgagctca  
240

aagagaagtt ccaacgcatg tgtgacaaat ccatgatcaa gagcagatac atgtatctaa  
300

cagaagagat tttgaaagaa aatcctagtc tttgtgaata catggcacct tcattggatg  
360

ctagacaaga catggtggtg gttgaggtac ctagacttgg gaaggaggct gcagtcaagg  
420

ctatcaaaga atgggggtcaa ccaaaatcta agattacaca tttgatcttt tgcaccacaa  
480

gtggtgtaga catgcctggt gctgattacc aactcacaaa actcttagga cttcgtccat  
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gaaggccctg caactatttt ggccattggc actgcaaadc cagcaaaccg tgttgatcag  
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agcacatadc ctgattttta cttcaaaadc actaacagtg agcataaggt tgagcttaaa  
240

gagaaatttc agcgcatgtg tgataaatct atgatcaaga gcagatacat gtatctaaca  
300

gaagagattt tgaaagaaaa tcctagtctt tgtgaatata tggcaccttc attggatgct  
360

aggcaagaca tgggtggtggt tgaggtacct agacttggga aggaggctgc agtgaaagct  
420

atcaaagaat ggggtcaacc aaagtcaaag attactcact taatcttttg caccacaagt  
480

ggtgtagaca tgccctggagc tgattaccaa ctcacaaaac tcttaggtct tcgccccatat



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120

gcaaccattt tggccattgg tactgcaaat ccaccaaacc gtgttgagcg gagcacatat  
180

cctgatttct acttcaaaat tacaaacagt gagcacaaga ctgagctcaa agagaagttc  
240

caacgcatgt gtgacaaatc catgatcaag agcagatata tgtatctaac agaagagatt  
300

ttgaaagaaa atcctagtct ttgtgaatac atggcacctt cattggatgc taggcaagac  
360

atgggtggtg ttgaggtacc tagacttggg aaggaggctg cagtgaagc tatcaaagaa  
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tgggggtcaac caaagtcaaa gattactcac ttaatctttt gcaccacaag tgggtgtagac  
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atgcctggag ctgattacca actcacaaaa ctcttaggtc ttgcccata tgtgaaaagg  
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120

caactatddd ggccattggg actgcaaate cagcaaatcg tgttgaccag agtacatate  
180

ctgatttcta cttcaaaate actaacagtg agcataaggt tgagcttaaa gagaaatttc  
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agcgcattgt tgataaatct atgatcaaga gcagatacat gtatctaaca gaagagattt  
300

tgaaagaaaa tcctagtctt tgtgaatata tggcaccttc attggatgct aggcaagaca  
360

tggtgggtgg tgaggtacct agacttggga aggaggctgc agtgaaagct atcaaagaat  
420

gggggtcaacc aaagtcaaag attactcact taatcttttg caccacaagt ggtgttgaca  
480

tgcttggtgc cgattaccaa ctcacaaaac tcttaggtct tgccccatat gtgaagaggt  
540

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aaggccctgc aactattttg gccattggta ctgcaaatcc agcaaactcg gttgaccaga  
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aagagatttt gaaagaaaat cctagtcttt gtgaatacat ggcaccttca ttggatgcta  
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ggcaagacat ggtgggtggtt gaggtaccta gacttgggaa ggaggctgca gtgaaagcta  
420  
tcaaagaatg ggggtcaacca aagtcaaaga ttactcactt aatcttttgc accacaagtg  
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ggccctgcaa ccattttggc cattggcact gcaaatccac caaacctgtg tgagcagagc  
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acatatcctg atttctactt caaaattaca aacagtgagc acaagactga gctcaaagag  
240  
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300  
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360  
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420  
aaagaatggg gtcaaccaa gtcaaagatt actcacttaa tcttttgcac cacaagtggg  
480  
gttgacatgc ctggtgctga ttaccaactc aaaaaactct taggtcttcg cccatatgtg  
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120

caactatddd ggccattggg actgcaaata cagcaaatcg tggtgaccag agtacatata  
180

ctgatttcta cttcaaaaata actaacagtg agcataaggt tgagcttaaa gagaaatttc  
240

agcgcatgtg tgataaatct atgatcaaga gcagatacat gtatctaaca gaagagattt  
300

tgaaagaaaa tcctagtctt tgtgaatata tggnaacctt attgnatgnt agncaagaca  
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390

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120

aggccctgca accatttttgg ccattggcac tgcaaattcca ccaaaccgtg ttgagcagag  
180

cacatatcct gatttctact tcaaaattac aaacagtgtg cacaagactg agtcaaaga  
240

gaagttccaa cgcattgtgtg acaaattccat gatcaagagc agatacatgt atctaacaga  
300

agagattttg aaagaaaatc ctagtctttg tgaatacatg gcacottcat tggatgctag  
360

acaagacatg gtgggtggtg aggtacctag acttggaag gaggtgcag tcaaggctat  
420

caaagaatgg ggtcaaccaa aatctaagat tacacatttg atcttttgca ccacaagtgg

480

tgtanacatg cctgggtgctg attaccnact ngcaaaactn ttaggacttn gcccatatgt  
540

gangaggcgc gtgntgmncc n  
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120  
ggccctgcaa ctattttggc cattgggtact gcaaattccag caaatcgtgt tgaccagagt  
180  
acatatcctg atttctactt caaaatcact aacagtgagc ataagggtga gcttaaagag  
240  
aaatttcagc gcatgtgtga taaatctatg atcaagagca gatacatgta tctaacagaa  
300  
gagattttga aagaaaatcc tagtctttgt gaatacatgg caccttcatt ggatgctagg  
360  
caagacatgg tgggtggtga ggtacctaga cttgggaagg aggctgcagt gaaagctatc  
420  
aaagaatggg gtcaaccaa gtcaaagatt actcacttaa tcttttgcac cacaagtggg  
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120

gccctgcaac tatitttgcc attggtactg caaatccagc aaaccgtgtt gatcagagta  
180

catatcctga tttctacttc aaaatcacta acagtgcagc taaagttgag ctcaaagaga  
240

aattccagcg catgtgtgat aaatctatga tcaagagcag atacatgtat ctaacagaag  
300

agattttgaa agaaaatcct agtctttgtg aatacatggc accttcattg gatgctaggc  
360

aagatatggg ggcgttgag gtacctanac ttgnaaagga ggctgcnnntg aaggctatta  
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120  
ccctgcaacc attttggcca ttggcactgc aaatccacca aaccgtgttg agcagagcac  
180  
atatcctgat ttctacttca aaattacaaa cagtgagcac aagactgagc tcaaagagaa  
240  
gttccaacgc atgtgtgaca aatccatgat caagagcaga tacatgtatc taacagaaga  
300  
gattttgaaa gaaaatccta gtctttgtga atacatggca ccttcattgg atgctaggca  
360  
agacatggtg gtggttgagg tacctagact tgggaaggag gctgcagtga aagctatcaa  
420  
agaatggggg caaccaaagt caaagattac tcacttaatc ttttgcacca caagtgggtg  
480  
agacatgcct ggagctgatt accaactcac aaaactctta ggtcttcgcc catatgtgaa  
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ctgcaaccat tttggccatt ggccactgcaa atccaccaaa ccgtgttgag cagagcacat  
180  
atcctgattt ctacttcaaa attacaaaca gtgagcacia gactgagctc aaagagaagt  
240  
tccaacgcat gtgtgacaaa tccatgatca agagcagata catgtatcta acagaagaga  
300  
ttttgaaaga aaatcctagt ctttgtgaat acatggcacc ttcattggat gctaggcaag  
360  
acatgggtggt gggtgaggta cctagacttg ggaaggaggc tgcagtcaag gccattaaag  
420  
aatgggggtca accaaagtca aagattactc acttaatctt ttgcaccaca agtgggtgttg  
480  
acatgcctgg tgctgattac caactcacia aactcttagg tcttcgcccc tatgtgaaaa  
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120

tgcaaccatt ttggccattg gcaactgcaaa tccaccaaac cgcgttgagc agagcacata  
180

tcctgatttc tacttcaaaa ttacaaacag tgagcacaag actgagctca aagagaagtt  
240

ccaacgcattg tgtgacaaat ccatgatcaa gagcagatac atgtatctaa cagaagagat  
300

tttgaaagaa aatcctagtc tttgtgaata catggcacct tcattggatg ctaggcaaga  
360

catgggtggtg gttgaggtac ctagacttgg gaaggaggct gcagtcaagg ctatcaaaga  
420

atgggggtcaa ccaaaatcta agattacaca tttgatcttt tgcaccacaa gtgggtgtaga  
480

catgcctggt gctgattacc aactcacaaa actcctggga cttcgtccat atgtgaagag  
540

attcatgatg taccaacaag gctgctttgc aggtgggacg gttcttcg  
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120

aaccattttg gccattggca ctgcaaatacc accaaaccgt gttgagcaga gcacatatcc

180

tgattttctac ttcaaaatta caaacagtga gcacaagact gagctcaaag agaagttcca  
240

acgcatgtgt gacaaatcca tgatcaagag cagatacatg tatctaacag aagagatttt  
300

gaaagaaaat cctagtcttt gtgaatacat ggcaccttca ttggatgcta ggcaagacat  
360

ggtgggtggtt gaggtaccta gacttgggaa ggaagctgca gtcaaggcca ttaaagaatg  
420

gggtcagcca aagtcaaaga ttactcactt aatcttttgc accacaagtg gtgttgacat  
480

gcctggagct gattaccaac tcacaaaact cttaggtctt cgcccatatg tgaaaaggta  
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tatgatgtac caacaagggtt gttttgcagg aggcacggtg cttcgtttgg caaaagattt  
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120

tttggccatt ggtactgcaa atccagcaaa ccgtgttgat cagagtacat atcctgattt  
180

ctacttcaaa atcactaaca gtgagcataa gggttgagctc aaggagaaat tccagcgcat  
240

gtgtgacaaa tctatgatca agagcagata catgtatcta acagaagaga ttttgaaaga  
300



aaatcctagt ctttgtgaat acatggcacc ttcattggat gctaggcaag atatggtggt  
360

cgttgaggta cctagactgg gaaaggaagc tgcagtgaag gctattaaag aatgggggtca  
420

accaaagtca aagattactc acttaatctt ttgcactaca agtgggtgtag acatgcctgg  
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120

ccattttggc cattggtact gcaaatccag caaaccgtgt tgatcagagt acatatcctg  
180

atttctactt caaaatcact aacagtgagc ataagggtga gctcaaggag aaattccagc  
240

gcatgtgtga taaatctatg atcaagagca gatacatgta tctaacagaa gagattttga  
300

aagaaaatcc tagtctgtgt gagtacatgg caccttcatt ggatgctagg caagacatgg  
360

ttgtgggtga ggtacctaga cttggaaagg aggctgcagt caaggccatt aaagaatggg  
420

gtcaacaaaa gtcaaagatt actcacttaa tcttttgcac cacaagtggg gtagacatgc  
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ctgggtgctga ttaccaactc acaaaactct tangtcttcg tccatacgtg aagagggaca  
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120

ccattttggc cattggcact gcaaattcac caaaccgtgt tgagcagagc acatatcctg  
180

atttctactt caaaattaca aacagtgagc acaagactga gctcaaagag aagttccaac  
240

gcatgtgtga caaatccatg atcaagagca gatacatgta tctaacagaa gagattttga

300

aagaaaatcc tagtctttgt gaatacatgg caccttcatt ggatgctagg caagacatgg  
360

tggtgggtga ggtacctaga cttgggaagg aggctgcagt caaggccatt aaagaatggg  
420

gtcaacaaaa gtcaaagatt actcacttaa tcttttgcac cacaagtggg gttgacatgc  
480

ctgggtgctga ttaccaactc acaaaactct taggtcttcg cccatatgtg aaaaggtata  
540

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&lt;211&gt; 546

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tggttactgt ttctgaaatt cgcaaggctc aaagggctga aggccctgca accatttttg  
120

ccattggtac tgcaaatcca gcaaaccgtg ttgatcagag tacatatoct gattttctact  
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tcaaaatcac taacagtga cataagggtg agctcaagga gaaattccag cgcattgtgtg  
240

ataaatctat gatcaagagc agatacatgt atctaacaga agagattttg aaagaaaatc  
300

ctagtctgtg tgagtacatg gcaccttcat tggatgctag gcaagacatg gttgtgggtg  
360

aggtagcctag acttggaaag gaggtctgcag tcaaggccat taaagaatgg ggtcaaccaa  
420

agtcaaagat tactcactta atcttttgca ccacaagtgg ttagacatg cctgggtgctg  
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ttgggtactgc aaatccagca aaccgtgttg atcaaagtac atatcctgat ttctacttca  
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aaatcactaa cagtgagcat aaggttgagc tcaaagagaa attccagcgc atgtgtgata  
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aatctatgat caagagcaga tacatgtatc caacagaaga gattttgaaa gaaaatccta  
300

gtctttgtga atacatggca ccttcattgg atgctagaca agacatgggtg gtgggtgagg  
360

tacctagact tggaaaggag gctgcagtga aggccattaa agaatgggggt caacaaaaat  
420

ctaagattac acatttgatc ttttgcacca caagtgggtg agacatgcct ggtgctgatt  
480

accagctcac aaaactctta ggtcttcgtc catatgtgaa aaggtatatg atgtaccaac  
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aagggtgctt tgcagggtggg acggtgcttc gtttggccaa gg  
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120

ttggccattg gtactgcaaa tccagcaaac cgtgttgatc aaagtacata tcttgatttc  
180

tacttcaaaa tctaatacag tgagcataag gttgagctca aagagaaatt ccagcgcnng  
240

tgtgataaat ctatgatcaa gagcagatac atgtatctaa cagaagagat tttgaaagaa  
300

aatcctagtc tttgtgaata catggcacct tcattggatg ctagacaaga catgggtggtg  
360

gttgaggtac ctagacttgg aaaggaggct gcagtgaagg ccattaaaga atgggggtcaa  
420

ccaaaatcta agattacaca tttgatcttt tgcaccacaa gtggtgtana catgcctggt  
480

gctgattacc agctcacaaa actcttaggt cttcgtccat atgtgaaaag ggatatgatg  
540

taccaacaag ggtgctttgc aggtgggacc gtgcttcntt tggccaanga tttggcccan  
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120

caaaccgtgt tgagcagagc acatatcctg atttctactt caaaattaca aacagtgagc  
180

acaagactga gctcaaagag aagttccaac gcatgtgtga caaatccatg atcaagagca  
240

67/390

gatacatgta tctaacagaa gagattttga aagaaaatcc tagtctttgt gaatacatgg  
300  
caccttcatt ggatgctagg caagacatgg tgggtggttga ggtacctaga cttgggaagg  
360  
aggctgcagt gaaagctatc aaagaatggg gtcaaccaaa gtcaaagatt actcacttaa  
420  
tcttttgcac cacaagtggg gtagacatgc ctggagctga ttaccaactc acaaaactct  
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120

gagcacatat cctgatttct acttcaaaat tacaaacagt gagcacaaga ctgagctcaa  
180

agagaagttc caacgcatgt gtgacaaatc catgatcaag agcagatata tgtatctaac  
240

agaagagatt ttgaaagaaa atcctagtct ttgtgaatac atggcacctt cattggatgc  
300

taggcaagac atggtggtgg ttgaggtacc tagacttggg aaggaggctg cagncnaggc  
360

cattaaanaa tggggncnac caaagncaaa gattactcac ttaatctttt gcaccacaag  
420

tggtgctgac atgnctggtg ctgattacca actcacaaaa ctcttaggnc ttcccccata  
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120  
cacatatcct gattttctact tcaaaattac aaacagttag cacaagactg agtcaaaga  
180  
gaagttccaa cgcattgtgtg acaaatacat gatcaagagc agatacatgt atctaacaga  
240  
agagattttg aaagaaaatc ctagtctttg tgaatacatg gcaccttcat tggatgctag  
300  
gcaagacatg gtggtggttg aggtacctag acttggaag gaggtgcag tcaaggccat  
360  
taaagaatgg ggttaaccaa agtgaaagat tactnactta atcttttgca ccacaagtgg  
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120  
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180  
gaagttccaa cgcattgtgtg acaaatacat gatcaagagc agatacatgt atctaacaga  
240  
agagattttg aaagaaaatc ctagtctttg tgaatacatg gcaccttcat tggatgctag  
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gcaagacatg gtggtggttg aggtacctag acttggaag gaggtgcag tgaaagctat  
360  
caaagaatgg ggttaaccaa agtcaaagat tactcactta atcttttgca ccacaagtgg  
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tgtagacatg cctggagctg attaccaact cacaaaactc ttaggtcttc gccatatgt  
480  
gaaaaggtat atgatgtacc aacaagggtg ttttgcagga ggcacggtgc ttcgtttggc  
540  
aaaagatttg gccgagaaca acaaagggtg tcgtgtgttg gttgtttgtt ctgaagtcac  
600

tgcagttaca ttccgtggcc cgagtgcacac tcacttggac agtcttgttg gacaagcatt  
660

gtttggagat ggagcagctg cacttatcgt tggttctgat ccagtgccag aaattgagaa  
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aggccctgca accatttttg ccattggcac tgcaaatcca ccaaaccgtg ttgagcagag  
120

cacatatcct gatttctact tcaaaattac aaacagttag cacaagactg agctcaaaga  
180

gaagttccaa cgcattgtgtg acaaatccat gatcaagagc agatacatgt atctaacaga  
240

agagattttg aaagaaaatc ctagtctttg tgaatacatg gcaccttcat tggatgctag  
300

acaagacatg gtggtgggtg aggtacctag acttgggaag gaggctgcag tcaaggctat  
360

caaagaatgg ggtcaaccaa aatctaagat tacacatttg atcttttgca ccacaagtgg  
420

tgtagacatg cctgggtgctg attaccaact cacaaaactc ttaggacttc gtccatatgt  
480

gaagaggtag atgatgtacc aacaagggtg ctttgcaggt ggggcggttc ttcgtttggc  
540

taaagatttg gccgagaaca acaaagggtgc tcgtgtgttg gttgtttggt ctgaagt  
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120

ctgagctcaa agagaagttc caacgcatgt gtgacaaatc catgatcaag agcagatata  
180

tgtatctaac agaagagatt ttgaaagaaa atcctagtct ttgtgaatac atggcacctt  
240

cattggatgc tagacaagac atggtggttg ttgaggtacc tagacttggg aaggaggccg  
300

cagtcaaggc tatcaaagaa tgggggtcaac caaaatctaa gattacacat ttgatctttt  
360

gcaccacaag tgggtgtagac atgcctggtg ctgattacca actcacaaaa ctcttaggac  
420

ttcgtccata tgtgaagagg tacatgatgt accaacaagg gtgctttgca ggtgggacgg  
480

ttcttcgttt ggctaaagat ttggccgaga acaacaaagg tgctcgtgtg ttggttgttt  
540

gttctgaagt aactgcagtc acattccgtg gcccgagtga cactcacttg  
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agaaaatttca gcgcatgtgt gataaatcta tgatcaagag cagatacatg tatctaacag  
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tcaaagaatg ggggtcaacca aagtcaaaga ttactcactt aatcttttgc accacaagtg  
360  
gtgttgacat gcttggtgct gattaccaac tcacaaaact cttaggtctt cgcccatatg  
420  
tgaagaggta catgatgtac caacaagggt gctttgcagg tgggacgggt cttcgtttgg  
480  
ccaaggattht ggccgagaac aacaaagggtg ctctgtgtgtt gggtgtttgc tctgaagtaa  
540  
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120  
aaatttcagc gcatgtgtga taaatctatg atcaagagca gatacatgta tctaacagaa  
180  
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240  
caagacatgg tgggtggtga ggtacctaga cttgggaagg aggctgcagt gaaagctatc  
300  
aaagaatggg gtcaaccaa gtcaaagatt actcacttaa tcttttgcac cacaagtgg  
360  
gttgacatgc ctggtgctga ttaccaactc aaaaactct taggtcttcg cccatatgtg  
420  
aagagggtaca tgatgtacca acaaggggtgc tttgcagggtg ggacgggtct tcgtttggcc  
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aaggatttgg ccgagaacaa caaagggtgct cgtgtgttgg ttgtttgctc tgaagtaacc  
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120  
caccttcatt ggatgctagg caagacatgg tgggtggtga ggtacctaga cttgggaagg  
180  
aggctgcagt gaaagctatc aaagaatggg gtcaacaaaa gtcaaagatt actcacttaa  
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360  
ggacgggttc tcgtttggcc aaggatttgg ccgagaacaa caaagggtgct cgtgtgttgg  
420  
ttgtttgttc tgaagtaacc gcagtcacat tccgcggccc cagtgcactc catttgaca  
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559

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420  
catttgaca gtcttggttg acaagcacta tttggagatg gagctgctgc tctcattgtt  
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120  
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300  
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gattgtatca aagaacatta ataaagcatt ggtcgaggct ttccaaccat taggaatttc  
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180  
gggcttaaaa cctgaaaaaa tgaaggccac cagagatgta cttagtgaat atggtaacat  
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480  
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540  
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Leu Asn Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His Pro Gly  
 35 40 45

Gly Pro Ala Ile Leu Asp Gln Val Glu Ile Lys Leu Gly Leu Lys Pro  
 50 55 60

Glu Lys Met Lys Ala Thr Arg Asp Val Leu Ser Glu Tyr Gly Asn Met  
 65 70 75 80

Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Gln Lys Lys Ser  
 85 90 95

Ala Glu Asn Gly Leu Lys Thr Thr Gly Glu Gly Leu Asp Trp Gly Val  
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Ser Val Ala Ile  
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180  
gggcttaaaa cctgaaaaaa tgaaggccac cagagatgta cttagtgaat atggtaacat  
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gtcaagtgca tgtgtattgt tcatcttaga tgagatgaga aagaaatcgg ctgaaaatgg  
300  
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taccattgaa actgttggtc tacatagtgt ggctatatga gaatgagaga cttgatttgt  
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ttaaataaat atatcgtaa tagctattat tttagtgtct gtttcttttt actaaactat  
300  
attttatttt agtatttgct attgatttga aataaatatt gtctctttaa ctgaaaaaaa  
360  
aaa

363

<210> 68  
<211> 363  
<212> DNA  
<213> *Trifolium repens*

<400> 68  
gagatgccta agaaatcggc tcaaaatgga ctgaaaacca caggagaagg acttgactgg  
60

ggtgtgttgt ttggatttgg accaggactt accattgaaa ctgttggtct acatagtgtg  
120

gctatatgag aatgcgagac ttgattgttt tgtattgtat tgtattgtat tgtattactt  
180

ttaatcttgc ttgaatttcc atttaacaat aaatatggag ttcaataagt accatcagtg  
240

ttaaaataat atatcgtaa tagctattat tttagtgtct gtttcttttt actaaactat  
300

atatttatttt agtatttgct attgatttga aataaatatt gtcctcttaa ctgaaaaaaaa  
360

aaa  
363

<210> 69  
<211> 897  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<222> (2)..(2)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (26)..(26)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (33)..(33)  
<223> Any nucleotide

<400> 69  
gnntcaatct gttgtgcata aaattncttt gcnatagaaa accatacaca tttgatcttg  
60

caaagaagaa atatgggaga cgaaggata gtgagagggtg tcacaaagca gacaaccct  
120

gggaaggcta ctatattggc tcttggcaag gcattccctc accaacttgt gatgcaagag  
180

tatttagttg atgggtattt tagggacact aattgtgaca atcctgaact taagcagaaa  
240

cttgctagac tttgtaagac aaccacggta aaaacaaggt atgttggttat gaatgaggag  
 300  
 atactaaaga aatatccaga acttggtgtc gaaggcgcct caactgtaaa acaacgttta  
 360  
 gagatatgta atgaggcagt aacacaaatg gcaattgaag cttcccaagt ttgcctaaag  
 420  
 aattggggta gatccttatc ggacataact catgtgggtt atgtttcatc tagtgaagct  
 480  
 agattacccg gtggtgacct atacttgtca aaaggactag gactaaaccc taaaattcaa  
 540  
 agaaccatgc tctatttctc tggatgctcg ggaggcgtag ccggccttcg cgttgcgaaa  
 600  
 gacatagctg agaacaaccc tggaagtaga gttttgcttg ctacttctga aactacaatt  
 660  
 attggattca agccaccaag tgttgataga ctttatgata ttggttggtgt ggcactcttt  
 720  
 ggagatggtg ctggtgctat gataattggc tcagacccaa tacttgaaac tgagactcca  
 780  
 ttgtttgagc ttcatacttc agctcaggag tttataccag acacagagaa gaaaatagat  
 840  
 gggcggctga cggaggaggg cataagtttc acgctagcga gggaactgcc gcagata  
 897

<210> 70  
 <211> 275  
 <212> PRT  
 <213> *Trifolium repens*

<400> 70

Met Gly Asp Glu Gly Ile Val Arg Gly Val Thr Lys Gln Thr Thr Pro  
 1 5 10 15

Gly Lys Ala Thr Ile Leu Ala Leu Gly Lys Ala Phe Pro His Gln Leu  
 20 25 30

Val Met Gln Glu Tyr Leu Val Asp Gly Tyr Phe Arg Asp Thr Asn Cys  
 35 40 45

Asp Asn Pro Glu Leu Lys Gln Lys Leu Ala Arg Leu Cys Lys Thr Thr  
 50 55 60

Thr Val Lys Thr Arg Tyr Val Val Met Asn Glu Glu Ile Leu Lys Lys  
 65 70 75 80

Tyr Pro Glu Leu Val Val Glu Gly Ala Ser Thr Val Lys Gln Arg Leu  
 85 90 95

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Glu Ile Cys Asn Glu Ala Val Thr Gln Met Ala Ile Glu Ala Ser Gln  
                   100                  105                  110  
 Val Cys Leu Lys Asn Trp Gly Arg Ser Leu Ser Asp Ile Thr His Val  
                   115                  120                  125  
 Val Tyr Val Ser Ser Ser Glu Ala Arg Leu Pro Gly Gly Asp Leu Tyr  
                   130                  135                  140  
 Leu Ser Lys Gly Leu Gly Leu Asn Pro Lys Ile Gln Arg Thr Met Leu  
                   145                  150                  155                  160  
 Tyr Phe Ser Gly Cys Ser Gly Gly Val Ala Gly Leu Arg Val Ala Lys  
                   165                  170                  175  
 Asp Ile Ala Glu Asn Asn Pro Gly Ser Arg Val Leu Leu Ala Thr Ser  
                   180                  185                  190  
 Glu Thr Thr Ile Ile Gly Phe Lys Pro Pro Ser Val Asp Arg Pro Tyr  
                   195                  200                  205  
 Asp Leu Val Gly Val Ala Leu Phe Gly Asp Gly Ala Gly Ala Met Ile  
                   210                  215                  220  
 Ile Gly Ser Asp Pro Ile Leu Glu Thr Glu Thr Pro Leu Phe Glu Leu  
                   225                  230                  235                  240  
 His Thr Ser Ala Gln Glu Phe Ile Pro Asp Thr Glu Lys Lys Ile Asp  
                   245                  250                  255  
 Gly Arg Leu Thr Glu Glu Gly Ile Ser Phe Thr Leu Ala Arg Glu Leu  
                   260                  265                  270  
 Pro Gln Ile  
                   275

<210> 71  
 <211> 577  
 <212> DNA  
 <213> Trifolium repens

<220>  
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 <223> Any nucleotide

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<220>



<221> misc\_feature  
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<220>  
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<400> 71  
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120  
gggaaggcta ctatattggc ttttggcaag gcattccctc accaacttgt gatgcaagag  
180  
tgtttagttg atgggtatct tagggacact aattgtgaca atcctgaact taagcagaaa  
240  
cttgctagac tttgtaagac aaccacggta aaaacaaggt atgttggtat gaatgaggag  
300  
atactaaaga aatatccaga acttggtgtc gaaggcgcct caactgtaa acaacgttta  
360  
gagatatgta atgaggcagt aacacaaatg gcaattgaag cttccaagt ttgcctaaag  
420  
aattggggta gatccttatc ggacataact catgtgggtt atgtttcatc tagtgaagct  
480  
agattacccg gtggtgacct atacttgtca aaaggactag gactaaaccc taaaattcaa  
540  
agaaccatgc tctatttctc tggatgctcg ggaggcg  
577

<210> 72  
<211> 599  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<222> (12)..(12)  
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<220>  
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<223> Any nucleotide

<220>  
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<222> (23)..(23)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (30)..(30)  
<223> Any nucleotide

<400> 72  
tcaatctgtt gngcattttt ttncttttgc atagaaaacc atacacattt gatcttgctt  
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agaagaaata tgggagacga aggtatagt agaggtgtca caaagcagac aacccttggg  
120  
aaggctacta tattggctct tggcaaggca ttccctcacc aacttgtgat gcaagagtgt  
180  
ttagttgatg gttatttttag ggacactaat tgtgacaatc ctgaacttaa gcagaaactt  
240  
gctagacttt gtaagacaac cacggtaaaa acaaggatatg ttgttatgaa tgaggagata  
300  
ctaaagaaat atccagaact tgttgtcgaa ggcgcctcaa ctgtaaaaca acgttttagag  
360  
atatgtaatg aggcagtaac acaaattggca attgaagctt cccaagtttg cctaaagaat  
420  
tggggtagat ccttatcgga cataactcat gtggttttatg tttcatctag tgaagctaga  
480  
ttacccggtg gtgacctata cttgtcaaaa ggactaggac taaaccctaa aattcaaaga  
540  
accatgctct atttctctgg atgctcggga ggcgtagccg gccttcgcgt tgcgaaaga  
599

<210> 73  
<211> 581  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<220>  
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<222> (26)..(26)  
<223> Any nucleotide

<400> 73  
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gaaatatggg agacgaaggt atagtgagag gtgtcacaaa gcagacaacc cctgggaagg

120

ctactatatt ggctcttggc aaggcattcc ctcaccaact tgtgatgcaa gagtatttag  
180

ttgatgggta ttttagggac actaattgtg acaatcctga acttaagcag aaacttgcta  
240

gactttgtaa gacaaccacg gtaaaaacaa ggtatgttgt tatgaatgag gagatactaa  
300

agaaatatcc agaacttggt gtcgaaggcg cctcaactgt aaaacaacgt ttagagatat  
360

gtaatgagggc agtaacacaa atggcaattg aagcttccca agtttgccta aagaattggg  
420

gtagatcctt atcggacata actcatgtgg tttatgtttc atctagttaa gctagattac  
480

ccggtgggtga cctatacttg tcaaaaggac taggactaaa ccctaaaatt caaagaacca  
540

tgctctatatt ctctggatgc tcgggaggcg tagccggcct t  
581

<210> 74  
<211> 588  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<222> (11)..(11)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (13)..(13)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (25)..(27)  
<223> Any nucleotides

<220>  
<221> misc\_feature  
<222> (36)..(36)  
<223> Any nucleotide

<400> 74  
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60

gaaatatggg agacgaagg atagtgaag gtgtcacaaa gcagacaacc cctgggaagg  
120

ctactatatt ggctcttggc aaggcattcc ctcaccaact tgtgatgcaa gagtggttag  
180

ttgatgggta ttttagggac actaattgtg acaatcctga acttaagcag aaacttgcta  
240

gactttgtaa gacaaccacg gtaaaaacaa ggtatgttgt tatgaatgag gagatactaa  
300

agaaatatcc agaacttggt gtcgaaggcg cctcaactgt aaaacaacgt ttagagatat  
360

gtaatgaggc agtaacacaa atggcaattg aagcttccca agtttgccca aagaattggg  
420

gtagatcctt atcggacata actcatgtgg tttatgtttc atctagtga gctagattac  
480

ccgggtggtga cctatacttg tcaaaaggac taggactaaa ccctaaaatt caaagaacca  
540

tgctctatct ctctggatgc tcgggaggcg tagccggcct tcgcgttg  
588

<210> 75

<211> 563

<212> DNA

<213> *Trifolium repens*

<400> 75

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60

cgaaggtata gtgagaggtg tcacaaggca gccaacccct gggaaggcta ctatattggc  
120

tcttggcaag gcattccctc accaacttgt gatgcaagag tatttagttg atggttatct  
180

tagggacact aattgtgaca atcctaagct taagcagaaa cttgctagac tttgcaagac  
240

aaccacagtg aaaacaaggt atgttggttat gaatgaagag atactaaaga aatatccaga  
300

acttactatc ggaggcacct cgacggtaaa acaacattta gagatatgta atgaggcagt  
360

aacacaaatg gcaattgaag cttcccaagt ttgcctaaag aattggggta gacccatctc  
420

agacataact catgtgggtt atgtttcatc tagtgaagct agattacctg gtggtgacct  
480

atacttgtca aaaggactag gactaaacct taaaattcaa agaaccatgc tctatttctc  
540

gggatgctcg ggaggcgtag ccg  
563

<210> 76

<211> 603

<212> DNA

<213> *Trifolium repens*

<220>

<221> misc\_feature

<222> (4)..(4)  
<223> Any nucleotide

<400> 76  
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60  
gtgagagggtg tcacaaagca gacaaccctt gggaaggcta ctatattggc tcttggcaag  
120  
gcattccctc accaacttgt gatgcaagag tatttagttg atgggtatct tagggacact  
180  
aattgtgaca atcctgacct taagcagaaa cttgctagac tttgtaagac aaccacggta  
240  
aaaacaagggt atgttggttat gaatgaggag atactaaaga aatatccaga acttggtgtc  
300  
gaaggcgcct caactgtaaa acaacgttta gagatatgta atgaggcagt aacacaaatg  
360  
gcaattgaag cttcccaagt ttgcctaaag aattggggta gatccttatt ggacataact  
420  
catgtggttt atgtttcatc tagtgaagct agattaccgc gtggtgacct atacttgtca  
480  
aaaggactag gactaaacct taaaattcaa agaaccatgc tctattttctc tggatgctcg  
540  
ggaggcgtag cgggccttcg cggtgcgaaa gacatagctg agaacaacct tggaagtaga  
600  
gtt  
603

<210> 77  
<211> 584  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (5)..(5)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (40)..(40)  
<223> Any nucleotide

<400> 77  
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60

aggcagtaac acaaatggca attgaagcct cccaagtttg cctaaagaat tggggtagac  
120

ccttatcaga cataactcat gtggtttatg tttcctctag tgaagctaga ttacccgggtg  
180

gtgacctata tttgtcaaaa ggactaggac taaatcctaa aattcaaaga accatgctct  
240

atttctctgg atgctcagga ggcgtagccg gccttcgcgt tgcgaaagac atagctgaga  
300

acaaccctgg aagtagagtt ttgcttgcta cttctgaaac tacaattatt ggattcaagc  
360

caccaagtgt tgatagacct tatgatcttg ttgggtgtggc actcttttga gatgggtgctg  
420

gtgctatgat aattggctca gacccaatac ttgaaactga gactccattg tttgagcttc  
480

atacttcagc tcaggagttt ataccagaca cagagaagaa aatagatggg cggctgacgg  
540

aggagggcat aagtttcacg ctagcgaggg aactgccgca gata  
584

<210> 78  
<211> 735  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (47)..(47)  
<223> Any nucleotide

<400> 78  
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60

ttgctaagtc atcaaccatt ccaattcctt aatataacct atcagtactc accatctttt  
120

cttccctcct gctaacttta tacttagaga agatggtgaa agttaatgag atccgccagg  
180

cacagagagc tgaaggccct gccactgtgt tggcaatcgg cactgcaact cctccaaact  
240

gtgttgatca gactacatac cccgactact acttccgcat cacaacacgt gagcacaaga  
300

cagagctcaa agaaaaattc cagcgcattgt gtgacaaatc tatgattaag aagagatata  
360

tgcatttgac agaagagatt ttgaaggaga atccaagttt atgtgagtac atggcacctt  
420

cattggatgc aagacaagac atgggtggttg tggaagtacc aaggctagga aaagaggcag  
480

caacaaaggc aatcaaggaa tgggggtcaac ctaagtccaa gattaccac ctcactcttt  
540

gcaccacaag tgggtgtggac atgcccggtg ccgactatca gcttacaaag ctttttaggcc  
600

ttcgtccgca tgtgaagcgt tacatgatgt accaacaagg ttgttttgct ggtggcacgg  
660

tgcttcgttt ggctaaagac ttggctgaaa acaacaaagg tgcccgtgta ttggtggttt  
720

gttcagagat aactg  
735

<210> 79  
<211> 194  
<212> PRT  
<213> *Trifolium repens*

<400> 79

Met Val Lys Val Asn Glu Ile Arg Gln Ala Gln Arg Ala Glu Gly Pro  
1 5 10 15

Ala Thr Val Leu Ala Ile Gly Thr Ala Thr Pro Pro Asn Cys Val Asp  
20 25 30

Gln Ser Thr Tyr Pro Asp Tyr Tyr Phe Arg Ile Thr Asn Ser Glu His  
35 40 45

Lys Thr Glu Leu Lys Glu Lys Phe Gln Arg Met Cys Asp Lys Ser Met  
50 55 60

Ile Lys Lys Arg Tyr Met His Leu Thr Glu Glu Ile Leu Lys Glu Asn  
65 70 75 80

Pro Ser Leu Cys Glu Tyr Met Ala Pro Ser Leu Asp Ala Arg Gln Asp  
85 90 95

Met Val Val Val Glu Val Pro Arg Leu Gly Lys Glu Ala Ala Thr Lys  
100 105 110

Ala Ile Lys Glu Trp Gly Gln Pro Lys Ser Lys Ile Thr His Leu Ile  
115 120 125

Phe Cys Thr Thr Ser Gly Val Asp Met Pro Gly Ala Asp Tyr Gln Leu  
130 135 140

Thr Lys Leu Leu Gly Leu Arg Pro His Val Lys Arg Tyr Met Met Tyr  
145 150 155 160

Gln Gln Gly Cys Phe Ala Gly Gly Thr Val Leu Arg Leu Ala Lys Asp  
165 170 175

Leu Ala Glu Asn Asn Lys Gly Ala Arg Val Leu Val Val Cys Ser Glu

Ile Thr

<210> 80  
<211> 574  
<212> DNA  
<213> *Trifolium repens*

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<222> (47)..(47)  
<223> Any nucleotide

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120  
cttcctccct gctaacttta gaatcagaga agatggtgaa tgттаатгag atccgccagg  
180  
cacagagagc tgaaggccct gccactgtgt tggcaatcgg cactgcaact cctccaaact  
240  
gtgtcgatca gactacatac ccggactact acttccgcat cacaacagt gagcacaaga  
300  
cagagctcaa agaaaaattc cagcgcgtgt gtgacaaatc tatgattaag aagagatata  
360  
tgcatttgac agaagagatt ttgaaggaga atccaagttt atgtgagtac atggcacctt  
420  
cattggatgc aagacaagac atgggtggttг tggaaгtacc aaggctagga aaagaggcag  
480  
caacaaaggc aattaaggaa tgggggtcaac ctaagtccaa gattaccacac ctcatctttt  
540  
gcaccaccag tgggtgtcgac atgcccgggtg ccgc  
574

<210> 81  
<211> 597  
<212> DNA  
<213> *Trifolium repens*

<400> 81  
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ttgctaagtc atcaaccatt ccaattcctt aatataacct atcagtactc accatctttt  
120  
cttcctccct gctaacttta gactcagaga agatggtgaa tgттаатгag atccgccagg  
180  
cacagagagc tgaaggccct gccaccgtgt tggcaatcgg cactgcaact cctccaaact



240

gtgtcgatca gaggatcatc ccagactact acttccgcat cacaacagc gagcacaaga  
300

cagagctcaa agaaaaattc cagcgcatgt gtgacaaatc tatgattaag aagagatata  
360

tgcatttgac agaagagatt ttgaaggaga atccaagttt atgtgagtag atggcacctt  
420

cattggatgc aagacaagac atgggtggtt tggaagtacc aaggctagga aaagaggctg  
480

caacaaaggc tatcaaggaa tgggggtcaac ctaagtccaa gattactcac ctcatctttt  
540

gcaccacaag tgggtgtggac atgcctggcg ccgactatca gcttacaag ctttttag  
597

&lt;210&gt; 82

&lt;211&gt; 616

&lt;212&gt; DNA

<213> *Trifolium repens*

&lt;220&gt;

&lt;221&gt; misc\_feature

&lt;222&gt; (70)..(70)

&lt;223&gt; Any nucleotide

&lt;220&gt;

&lt;221&gt; misc\_feature

&lt;222&gt; (616)..(616)

&lt;223&gt; Any nucleotide

&lt;400&gt; 82

gtagcaacac acactttgat ttctttttga gtccttgcta cgtggcttta ccaaaaaacg  
60

ttgctaagtn atcaaccatt ccaattcctt aatataacct atcagtactc accatctttt  
120

cttcctccct gctaacttta gactcagaga agatggtgaa tgtaatgag atccgccagg  
180

cacagagagc tgaaggccct gccaccgtgt tggcaatcgg cactgcaact cctccaaact  
240

gtgtcgatca gaggatcatc ccagactact acttccgcat cacaacagc gagcacaaga  
300

cagagctcaa agaaaaattc cagcgcatgt gtgacaaatc tatgattaag aagagatata  
360

tgcatttgac agaagagatt ttgaaggaga atccaagttt atgtgagtag atggcacctt  
420

cattggatgc aagacaagac atgggtggtt tggaagtacc aaggctagga aaagaggctg  
480

caacaaaggc tatcaaggaa tgggggtcaac ctaagtccaa gattactcac ctcatctttt  
540

gcaccacaag tgggtgtggac atgcctggcg ccgactatca gcttacaaag cttttaggcc  
600

ttcgtccgca tgtgan  
616

<210> 83  
<211> 585  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (1)..(3)  
<223> Any nucleotides

<220>  
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<222> (7)..(7)  
<223> Any nucleotide

<220>  
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<220>  
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<222> (37)..(37)  
<223> Any nucleotide

<220>  
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<222> (61)..(61)  
<223> Any nucleotide

<220>  
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<222> (451)..(451)  
<223> Any nucleotide

<220>  
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<222> (507)..(507)  
<223> Any nucleotide

<400> 83  
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ntcaaccatt ccaattcctt aatataacct atcagtactc accatctttt cttcctccct  
120

gctaacttta gactcagaga agatggtgaa tgttaatgag atccgccagg cacagagagc  
180

tgaaggccct gccaccgtgt tggcaatcgg cactgcaact cctccaaact gtgttgatca  
240

gagtacatac cggactact acttccgcat cacaaacagt gagcacaaga cagagctcaa  
300

agaaaaattc cagcgcatgt gtgacaaatc tatgattaag aagagataca tgcatttaac  
360

agaagagatt ttgaaggaga atccaagttt atgtgagtac atggcacctt cattggatgc  
420

aagacaagac atggtgggtg tggaagtacc naggctagga aaagaggcag taacaaaggc  
480

aattagggaa tgggggtcaac ctaagtncaa gattaccac ctcattttt gcaccaccag  
540

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120

cctaaaaact gtgttgatca gagtacttac cccgactact atttccgaat cacaaacagc  
180

gaacacaaga ctgaactcaa agaaaaattc cagcgcatgt gtgacaaatc tatgattaag  
240

aagagataca tgcatttgac agaagagatt ttgaaggaga attcaagttt atgtgagtac  
300

atggcacctt cattggatgc aagacaagac atggtgggtg tggaagtacc aaggctagga  
360

aaagaggctg caacaaaggc tatcaaggaa tgggggtcaac ctaagtccaa gattactcac  
420

ctcatctttt gcaccacaag tgggtgtggac atgcctgggtg ccgactatca gcttacaag

480

cttttaggcc ttcgccgca tgtgaagcgt tatatgatgt accaacaagg ttgttttgct  
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120

cctaaaaact gtgttgatca gagtacttac cccgactact atttccgaat cacaaacagc  
180

gaacacaaga ctgaactcaa agaaaaattc cagcgcacgt gtgacaaatc tatgattaag  
240

aagagataca tgcatttgac agaagagatt ttgaaggaga attcaagttt atgtgagtac  
300

atggcacctt cattggatgc aagacaagac atggtggttg tggaagtacc aaggctagga  
360

aaagaggctg caacaaaggc tatcaaggaa tgggggtcaac ctaagtccaa gattactcac  
420

ctcatctttt gcaccacaag tgggtgtggac atgcctgggtg ccgactatca gcttacaaag  
480

cttttaggcc ttcgtccgca tgtgaagcgt tatatgatgt accaacaagg ttgttttgc  
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ttggtggttt gttcagag  
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120

aaaactgtgt tgatcagagt acttaccocg actactatct ccgaatcaca aacagcgaac  
180

acaagactga actcaaagaa aaattccagc gcatgtgtga caaatctatg attaagaaga  
240

gatacatgca tttgacagaa gagattttga aggagaattc aagtttatgt gagtacatgg  
300

caccttcatt ggatgcaaga caagacatgg tggttgtgga agtaccaagg ctaggaaaag  
360

aggctgcaac aaaggctatc aaggaatggg gtcaacctaa gtccaagatt actcacctca  
420

tcttttgcac cacaagtggg gtggacatgc ctggtgccga ctatcagctt acaaagcttt  
480

taggccttcg tccgcatgtg aagcgttata tgatgtacca acaagggtgt tttgctgggt  
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gcacgggtgct tcgtttggct aaagacttgg ctgaaaacaa caaagggtgcc cgtgtattgg  
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120  
tcagagtaca taccgggact actacttccg catcacaaac agtgagcaca agacagagct  
180  
caaagaaaaa ttccagcgca tgtgtgacaa atctatgatt aagaagagat acatgcattt  
240  
gacagangag attttgaagg agaatccaag tttatgtgag tacatggcac cttcattgga  
300  
tgcaagacaa gacatgggtg ttgtggaagt accaaggcta ggaaaagagg cagcaacaaa  
360  
ggcaattaag gaatggggcc aacctaagtc caagattacc cacctcatct tttgcaccac  
420  
cagtgggtgc gacatgcccg gtgccgacta tcagctcaca aagctcttag gcctgcgtcc  
480  
atatgtgaag cgttacatga tgtatcaaca aggttggttt gctgggtggca cggtgcttcg  
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571

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120  
gtacataccc ggactactac ttccgcatca caaacagtga gcacaagaca gagctcaaag  
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aaaaattcca gcgcatgtgt gacaaatcta tgattaagaa gagatacatg catttgacag  
240  
aagagatttt gaaggagaat ccaagtttat gtgagtacat ggcacottca ttggatgcaa  
300  
gacaagacat ggtggttgtg gaagtaccaa ggctaggaaa agaggcagca acaaaggcaa  
360  
ttaaggaatg gggccaacct aagtccaaga ttaccacact catcttttgc accaccagt  
420  
gtgtcgacat gcccggtgcc gactatcagc tcacaaagct cttaggcctg cgtccatattg  
480  
tgaagcgta catgatgtat caacaagggt gttttgctgg tggcacggtg cttcgtttgg  
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603

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120

cttaccgccga ctactatttc cgaatcacaa acagcgaaca caagactgag ctcaaagaaa  
180

aattccagcg catgtgtgac aaatctatga ttaagaagag atacatgcat ttgacagaag  
240

agatthttgaa ggagaatcca agthttatgtg agtacatggc accttcattg gatgcaagac  
300

aagacatggt ggtagtggaa gtaccaaggc taggaaaaga ggcagcaaca aaggcaatta  
360

aagaatgggg tcaacctaa g tccaagatta cccacctcat cthtttgcacc accagtgggtg  
420

tggacatgcc cggtgccgac tatcagctca caaagctctt aggcctgcgt ccatatgtga  
480

agcgttacat gatgtatcaa caaggthgtt ttgctgggtg cacggtactt cgtthtggtta  
540

aagacttggc tgaaaacaac aaaggthgccc gtgtgttggt ggthttgtt  
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cacaagactg agctcaaaga aaaattccag cgcattgtgtg acaaattctat gattaagaag  
120

agatacatgc atttgacaga agagattttg aaggagaatc caagnttatg tgagtacatg

180

gcaccttnat tggatgcaag acaagacatg gnggccgcc accnnccntc cncncaccn  
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caaccnccn  
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120

ttcctccctg ctaacttttag actcagtaga agatgggtgaa tgттаатgag atccgccagg  
180

cacagagagc tgaaggccct gccaccgtgt tggcaatcgg cactgcaact cctccaaact  
240

gtgttgatca gagtacatac ccggactact acttccgcat cacaacacgt gagcacaaga  
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cagagctcaa agaaaaattc cagcgcacgt gtaagatatt tatcttatac tccatgcatg  
360

tttttttctg ctgactgccg tgtttatata ttgttttggt ttgttcctta aatttgttat  
420

gtcactctca catgtacaaa acacttaaga ctaaactgca tatcattttt ttcagggaca  
480

aatctatgat taagaagaga tacatgcatt tgacagaaga gattttgaag gagaatccaa  
540

gtttatgtga gnacatggca ccttcttggg atgcaagaca agt  
583

<210> 92  
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<212> PRT  
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<400> 92

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Ala Thr Val Leu Ala Ile Gly Thr Ala Thr Pro Pro Asn Cys Val Asp  
 20 25 30

Gln Ser Thr Tyr Pro Asp Tyr Tyr Phe Arg Ile Thr Asn Ser Glu His  
 35 40 45

Lys Thr Glu Leu Lys Glu Lys Phe Gln Arg Thr Arg Asp Lys Ser Met  
 50 55 60

Ile Lys Lys Arg Tyr Met His Leu Thr Glu Glu Ile Leu Lys Glu Asn  
 65 70 75 80

Pro Ser Leu Cys Glu Xaa Met Ala Pro Ser Trp Asp Ala Arg Gln  
 85 90 95

<210> 93  
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 120

ttcctccctg ctaacttttag actcagagaa gatgggtgaat gttaatgaga tccgccaggc  
 180

acagagagct gaaggccctg ccaccgtgtt ggcaatcggc actgcaactc ctccaaactg  
 240

tgttgatcag agtacatacc cggactacta cttccgcata acaaacagtg agcacaagac

300

agagctcaaa gaaaaattcc agcgcacgtg taagatattt atcttatact ccatgcatgt  
360

ctttttctgc tgactgccgt gtttatatat tgttttgttt tgttccttaa atttggtatg  
420

tcactctcac atgtacaaaa cacttaagac taaactgcat atcatttttt tcagggacaa  
480

atctatgatt aagaagagat acatgcattt gacagaagag attttgaagg agaatccaag  
540

tttatgtgag nacatggcac cttcttgga tgcaagacaa gt  
582

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aacctatcag tactcaccat cttttcttcc tccttgctaa ctttagactc agtagaagat  
120

ggtgaatggt aatgagatcc gccaggcaca gagagctgaa ggccctg  
167

<210> 95  
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&lt;223&gt; Any nucleotide

&lt;400&gt; 95

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120acgtgctcct actcagggaa aggcaacgat acttgcatta ggaaaggctt tccccgccca  
180agtcctccct caagagtgcg tggtggaagg attcattcgc gacactaagt gtgacgatac  
240ttatattaag gagaaattgg agcgtctttg caaaaacaca actgtaaaaa caagatacac  
300agtaatgtca aaggagatct tagacaacta tccagagcta gccatagatg gaacaccaac  
360aataaggcaa aagcttgaaa tagcaaatcc agcagtagtt gaaatggcaa caagagcaag  
420caaagattgc atcaaagaat ggggaagggtc acctcaagat atcacacaca tagtctatgt  
480ttcctcgagc gaaattcgtc taccgggtgg tgacctttat cttgcaaatg aactcggctt  
540aaacagcgat gttaatcgcg taatgctcta tttcctcggt tgctacggcg gtgtcactgg  
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613

&lt;210&gt; 96

&lt;211&gt; 182

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 96

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20 25 30Lys Ala Phe Pro Ala Gln Val Leu Pro Gln Glu Cys Leu Val Glu Gly  
35 40 45Phe Ile Arg Asp Thr Lys Cys Asp Asp Thr Tyr Ile Lys Glu Lys Leu  
50 55 60Glu Arg Leu Cys Lys Asn Thr Thr Val Lys Thr Arg Tyr Thr Val Met  
65 70 75 80

Ser Lys Glu Ile Leu Asp Asn Tyr Pro Glu Leu Ala Ile Asp Gly Thr

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	85		90		95
Pro Thr Ile Arg Gln Lys Leu Glu Ile Ala Asn Pro Ala Val Val Glu	100		105		110
Met Ala Thr Arg Ala Ser Lys Asp Cys Ile Lys Glu Trp Gly Arg Ser	115		120		125
Pro Gln Asp Ile Thr His Ile Val Tyr Val Ser Ser Ser Glu Ile Arg	130		135		140
Leu Pro Gly Gly Asp Leu Tyr Leu Ala Asn Glu Leu Gly Leu Asn Ser	145		150		155
Asp Val Asn Arg Val Met Leu Tyr Phe Leu Gly Cys Tyr Gly Gly Val	165		170		175
Thr Gly Leu Arg Val Ala	180				

<210> 97  
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 120  
 acgtgctcct actcagggaa aggcaacgat acttgcatta ggaaaggctt tccccgccca  
 180  
 agtcctccct caagagtgc tgggtggaagg attcattcgc gacactaagt gtgacgatac  
 240  
 ttatattaag gagaaattgg agcgtctttg caaaaacaca actgtaaaaa caagatacac  
 300  
 agtaatgtca aaggagatct tagacaacta tccagagcta gccatagatg gaacaccaac  
 360  
 aataaggcaa aagcttgaaa tagcaaatcc agcagtagtt gaaatggcaa caagagcaag  
 420

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caaagattgc atcaaagaat ggggaaggtc acctcaagat atcacacaca tagtctatgt  
480

ttcctcgagc gaaattcgtc tacccggttg tgacctttat cttgcaaata aactcggctt  
540

aaacagcgat gttaatcgcg taatgctcta tttcctcggg tgctacggcg gtgtcactgg  
600

cttacgtgtc gcc  
613

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120

tgctcctact cagggaagg caacgatact tgcattagga aaggctttcc ccgcccaggt  
180

cctccctcaa gagggttg tggaaggatt cattcgcgac actaagtgtg acgatactta  
240

tattaaggag aaattggagc gtctttgcaa aaacacaact gtgaaaacaa gatacacagt  
300

aatgtcaaag gagatcttag acaactatcc agagctagcc atagatggaa caccaacaat  
360

aaggcaaaag cttgaaatag caaatccagc agtagttgaa atggcaacaa gagcaagcaa  
420

agattgcac aaagaatggg gaaggtcacc tcaagatata acacacatag tctatgtttc  
480

ctcgagcgaa attcgtctac ccggtggtga cctttatctt gcaaatgaac tcggcttaaa  
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120  
tactcagggg aaggcaacga tacttgcatt aggaaaggct ttccccgccc aggtcctccc  
180  
tcaagagtgc ttggtggaag gattcattcg cgacactaag tgtgacgata cttatattaa  
240  
ggagaaattg gagcgtcttt gcaaaaacac aactgtgaaa acaagatata cagtaatgtc  
300  
aaaggagatc ttagacaact atccagagct agccatagat ggaacaccaa caataaggca  
360  
aaagcttgaa atagcaaata cagcagtagt tgaaatggca acaagagcaa gcaaagattg  
420  
catcaaagaa tggggaaggt cacctcaaga tatcacacac atagtctatg tttcctcgag  
480  
cgaaattcgt ctaccgggtg gtgaccttta tcttgcaaat gaactcggct taaacagcga  
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575

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120  
aaaggcaacg atacttgcatt taggaaaggc tttccccgcc caagtcctcc ctcaagagtg  
180  
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240  
ggagcgtctt tgcaaaaaca caactgtaaa aacaagatac acagtaatgt caaaggagat  
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cttagacaac tatccagagc tagccataga tggaacacca acaataaggc aaaagcttga  
360  
aatagcaaat ccagcagtag ttgaaatggc aacaagagca agcaaagatt gcatcaaaga  
420  
atggggaagg tcacctcaag atatcacaca catagtctat gtttcctcga gcgaaattcg  
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540

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573

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120

ggacccagtg acactcacct tgatagcctt gtggggcaag cattgtttgg agatgggtgca  
180

gcagctgtga ttgttggttc agaccctttg ccagaagttg agaagccttt gtttgaattg  
240

gtatggaccg cacaacaat cgctccagat agtgaaggag ccattgatgg tcacottcgc  
300

gaagcagggc tgacattcca tctcctcaag gatgttccta gccttgtctc aaataacatt  
360

gagaaagcgc ttgttgatgc ctttcaacct ttgaatattt ctgactacaa ttccatcttt  
420

tggattgcac acccaggcgg accagcaatt cttgaccaag ttgaagctaa gttaggctta  
480

aagccagaga aaatgcaagc cactcgacat gtacttagcg aatatggtaa catgtcaagt  
540

gcgtgtgtgt tatttatctt ggatgagatg aggaggaagt caaaagaaga cggacttgcc  
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acaacag  
607

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<400> 102

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Asp Leu Ala Glu Asn Asn Lys Gly Ala Arg Val Leu Val Val Cys Ser  
20 25 30

Glu Ile Thr Ala Val Thr Phe Arg Gly Pro Ser Asp Thr His Leu Asp  
35 40 45

Ser Leu Val Gly Gln Ala Leu Phe Gly Asp Gly Ala Ala Ala Val Ile  
50 55 60

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Val Gly Ser Asp Pro Leu Pro Glu Val Glu Lys Pro Leu Phe Glu Leu
65                               70               75               80

Val Trp Thr Ala Gln Thr Ile Ala Pro Asp Ser Glu Gly Ala Ile Asp
85                               90               95

Gly His Leu Arg Glu Ala Gly Leu Thr Phe His Leu Leu Lys Asp Val
100                             105               110

Pro Ser Leu Val Ser Asn Asn Ile Glu Lys Ala Leu Val Asp Ala Phe
115                             120               125

Gln Pro Leu Asn Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His
130                             135               140

Pro Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Ala Lys Leu Gly Leu
145                             150               155               160

Lys Pro Glu Lys Met Gln Ala Thr Arg His Val Leu Ser Glu Tyr Gly
165                             170               175

Asn Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Arg
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Lys Ser Lys Glu Asp Gly Leu Ala Thr Thr
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120

ggacccagtg acatcatct tgatagcctt gtggggcaag cattgtttgg agatggtgca
180

gcagctgtga ttgttggttc agaccctttg ccagaagttg agaagccttt gtttgaattg
240

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gtatggacCG cacaacaat cgctccagat agtgaaggag ccattgatgg tcaccttcgc  
300

gaagcagggc tgacattcca tctcctcaag gatgttccta gccttgtctc aaataacatt  
360

gagaaagctc ttgttgatgc ctttcaacct ttgaatatct ccgattacaa ttccatcttt  
420

tggattgcac acccaggcgg accagcaatt cttgaccaag ttgaagctaa gttaggctta  
480

aagccagaga aaatgcaatc cactcgacat gtacttagcg aatatggtaa catgtcgagt  
540

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acaacag  
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cagtgcact caccttgata gccttgtggg gcaagcattg tttggagatg gtgcagcagc  
180

tgtgattgtt ggttcagacc ctttgccaga agttgagaag cttttgtttg aattggatatg  
240

gaccgcacaa acaatcgctc cagatagtga aggagccatt gatggtcacc ttgcggaagc  
300

agggctgaca ttccatctcc tcaaggatgt tcttagcctt gtctcaaata acattgagaa  
360

agcgcttggt gatgcctttc aacctttgaa tattttctgac tacaattcca tcttttggat

420

tgcacaccca ggcggaccag caattcttga ccaagttgaa gctaagttag gcttaaagcc  
480

agagaaaatg caagccactc gacatgtact tagcgaatat ggtaacatgt caagtgcgtg  
540

tgtgttatatt atcttggatg anatgaggag gaagtcaaaa gaacacngnc t  
591

&lt;210&gt; 105

&lt;211&gt; 590

&lt;212&gt; DNA

<213> *Trifolium repens*

&lt;220&gt;

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&lt;223&gt; Any nucleotide

&lt;400&gt; 105

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120

gacactcacc ttgatagcct tgtggggcaa gcattgtttg gagatgggtgc agcagctgtg  
180

attgttggtt cagacccttt gccagaagtt gagaagcctt tgtttgaatt ggtatggacc  
240

gcacaaacaa togtccaga tagtgaagga gccattgatg gtcaccttcg cgaagcaggg  
300

ctgacattcc atctcctcaa ggatgttcct agccttgtct caaataacat tgagaaagcg  
360

cttgttgatg cctttcaacc tttgaatatt tctgactaca attccatctt ttggattgca  
420

cacccaggcg gaccagcaat tcttgaccaa gttgaagcta agttaggctt aaagccagag  
480

aaaatgcaag ccactcgaca tgtacttagc gaatatggta acatgtcaag tgcgtgtgtg  
540

ttatttatct tggatgagat gaggaggaag tcaaaagaag acggacttgc  
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&lt;210&gt; 106

&lt;211&gt; 510

&lt;212&gt; DNA

<213> *Trifolium repens*

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120  
cccaacgttc aaatggccct gccactatct tagcttttgg cacagccact ccttctaact  
180  
gtgtcactca agctgattat cctgattact actttcgtat caccaacagc gaacatatga  
240  
ctgatcttaa ggaaaaattc aagcggatgt gtgatagatc aatgataaag aaacgttaca  
300  
tgcacctaac agaagacttt ctgaaggaga atccaaatat gtgtgaatac atggcaccat  
360

cactagatgt aagacgagac atagtgggtg ttgaagnacc aaagctaggt aaagaancac  
420

caaaaaaagc catatgngaa tggggacaac caaaatcnaa aatcacacat gcttggtttc  
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tgaccacttc cggtgntgac atgcccgggg  
510

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Ala Thr Ile Leu Ala Phe Gly Thr Ala Thr Pro Ser Asn Cys Val Thr  
20 25 30

Gln Ala Asp Tyr Pro Asp Tyr Tyr Phe Arg Ile Thr Asn Ser Glu His  
35 40 45

Met Thr Asp Leu Lys Glu Lys Phe Lys Arg Met Cys Asp Arg Ser Met  
50 55 60

Ile Lys Lys Arg Tyr Met His Leu Thr Glu Asp Phe Leu Lys Glu Asn  
65 70 75 80

Pro Asn Met Cys Glu Tyr Met Ala Pro Ser Leu Asp Val Arg Arg Asp  
85 90 95

Ile Val Val Val Glu Xaa Pro Lys Leu Gly Lys Glu Pro Lys Lys Ala  
100 105 110

Ile Xaa Glu Trp Gly Gln Pro Lys Xaa Lys Ile Thr His Ala Trp Phe  
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Leu Thr Thr Ser Gly Asp Met Pro Gly  
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120

atggctcana acaagctctt ggtgaagggt tgaaagaagc anttgaactt ggtcttgctca  
180

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Lys Asp Thr Lys Asp Ala Ile Val Glu Ala Ile Lys Gln Gly Tyr Arg  
 20 25 30

His Phe Asp Thr Ala Ala Ala Tyr Gly Ser Xaa Gln Ala Leu Gly Glu  
 35 40 45

Gly Leu Lys Glu Ala Xaa Glu Leu Gly Leu Val Thr Xaa Glu Glu Xaa  
 50 55 60

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Phe Val Thr Ser Lys Leu Trp Xaa Thr Glu Asn His Xaa Asn Leu  
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 120  
 gtccatcaaa tgatgttctt gcttcaatct ttgttgatgc aattgaagtt gggtatcgcc  
 180  
 atttcgattc tgcttctgtg tatggaacag aggaagccat aggaattgct ttagcaaaaag  
 240  
 ctttagaaaa agggcttatt aagagtagag atgaagtttt catcacttca aagccatgga  
 300  
 atactgatgc agattatgaa cttattgttc cagctctcaa gaccacattg aaaaagctgg  
 360  
 ggaaggagta tgtggatctt tatctgatcc attggccagt gagacttaga catgatcttg  
 420  
 aaaaccctgt tgttttcacc aaagaagatt tacttccctt tgatatagaa gggacatgga  
 480  
 aagctatgga agaatgttat aagttaggct tagcaaagtc tattggtata tgcaattatg  
 540  
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 600  
 tgga  
 604

<210> 111  
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<400> 111

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 1 5 10 15

Met Pro Val Ile Gly Met Gly Thr Ser Val Asp Asn Arg Pro Ser Asn  
 20 25 30

Asp Val Leu Ala Ser Ile Phe Val Asp Ala Ile Glu Val Gly Tyr Arg

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35

40

45

His Phe Asp Ser Ala Ser Val Tyr Gly Thr Glu Glu Ala Ile Gly Ile  
 50 55 60

Ala Leu Ala Lys Ala Leu Glu Lys Gly Leu Ile Lys Ser Arg Asp Glu  
 65 70 75 80

Val Phe Ile Thr Ser Lys Pro Trp Asn Thr Asp Ala Asp Tyr Glu Leu  
 85 90 95

Ile Val Pro Ala Leu Lys Thr Thr Leu Lys Lys Leu Gly Thr Glu Tyr  
 100 105 110

Val Asp Leu Tyr Leu Ile His Trp Pro Val Arg Leu Arg His Asp Leu  
 115 120 125

Glu Asn Pro Val Val Phe Thr Lys Glu Asp Leu Leu Pro Phe Asp Ile  
 130 135 140

Glu Gly Thr Trp Lys Ala Met Glu Glu Cys Tyr Lys Leu Gly Leu Ala  
 145 150 155 160

Lys Ser Ile Gly Ile Cys Asn Tyr Gly Thr Lys Lys Leu Thr Lys Leu  
 165 170 175

Leu Glu Thr Ala Thr Ile Thr Pro Ala Val Asn Gln Val  
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<210> 112  
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120

gtccatcaaa tgatgttctt gcttcaatct ttgttgatgc aattgaagtt gggtatcgcc  
180

atttcgattc tgcttctgtg tatggaacag aggaagccat aggaattgct ttagcaaaag  
240

ctttanaaaa agggcttatt aagagtanag atgaagnttt natcacttgc aagncatgga  
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120  
ccatcaaatg atgttcttgc ttcaatcttt gttgatgcaa ttgaagttgg ttatogccat  
180  
ttcgattctg cttctgtgta tggaacagag gaagccatag gaattgcttt agcaaaagct  
240  
ttagaaaaag ggcttattaa gagtagagat gaagttttca tcacttcaaa gccatggaat  
300  
actgatgcag attatgaact tattgttcca gctctcaaga ccacattgaa aaagctgggg  
360  
acggagtatg tggatcttta tctgatccat tggccagtga gacttagaca tgatcttgaa  
420  
aaccctgttg ttttcaccaa agaagattta cttccctttg atatagaagg gacatggaaa  
480  
gctatggaag aatgttataa gttaggctta gcaaagtcta ttggtatatg caattatggt  
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atgatgttct tgcttcaatc tttgttgatg caattgaagt tggttatcgc catttcgatt  
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ctgcttctgt gtatggaaca gaggaagcca taggaattgc tttagcaaaa gctttagaaa  
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aagggccttat taagagtaga gatgaagttt tcatcacttc aaagccatgg aatactgatg  
300  
cagattatga acttattggt ccagctctca agaccacatt gaaaaagctg gggacggagt  
360  
atgtggatct ttatctgac cattggccag tgagacttag acatgatctt gaaaaccctg  
420  
ttgtttttcac caaagaagat ttacttccct ttgatataga agggacatgg aaagctatgg  
480  
aagaatgtta taagttaggc ttagcaaagt ctattggtat atgcaattat ggtacaaaaa  
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240  
aagggttat taagagtaga gatgaagttt tcatcacttc aaagccatgg aatactgatg  
300  
cagattatga acttattgtt ccagctctca agaccacatt gaaaaagctg gggacggagt  
360  
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aagctatagg aattgcttta gcaaaagctt tagaaaaagg gcttattaag agtagagatg  
240  
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334

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 cactagagaa gagctttttg ttacttctaa actttgggtc actgaaaatc atcctcacct  
 420  
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 480  
 tttgatccat tggccactta gttctcagcc tggaaagttt tcattttccaa ttgatgtggc  
 540  
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480  
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360

129/390

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acttggactt gtatttgatc cactggccac ttagttctca gcccgaaaag ttttcatttc  
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caattgaggt ggcagatctc ttgccatttg atgtgagggg tgtttggcaa tccatggaag  
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180

aagaaagata caaaagatgc aatcattgaa gccatcaaac aaggttatag acactttgat  
240

attgctgctg cttatggctc agaacaagct cttggtgaag gtttgaaaga agcaattgaa  
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cttggctcttg tcactagaga agaccttttt gttacttcta aactttgggt cactgaaaat  
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catcctcacc ttgttattcc tgctcttcaa aaatctctca agactcttca attggagtac  
420

ttggacttgt atttgatcca ctggccactt agttctcagc ccggaaagtt ttcatttcca  
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attgaggtgg cagatctctt gccatttgat gtgaggggtg tttggcaatc catggaagaa  
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agaaagacac aaaagatgca atcattgaag ccatcaaaca aggntataga cactttgaaa  
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240  
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300  
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360  
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420  
tggacttgta ttgatccat tggccactta attctcancc tggaaagttt tcatttccca  
480  
ttgangtggc aaatctcttg ccatttnatg tgaanggtgt ttgggaatcc atggaaaaan  
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agacacaaaa gatgcaatca ttgaagccat caaacaaggt tatagacact ttgatactgc  
240

tgctgcttat ggctcagaac aagctcttgg tggaggtttg aaagaagcaa ttgaacttgg  
300

ccttgctact agagaagagc tttttgttac ttctaaactt tgggtcactg aaaatcatcc  
360

tcaccttggt gttcctgctc ttcaaaaatc tctcaagact cttcaattgg agtacttgga  
420

cttgatattg atccattggc cacttagttc tcagcctgga aagttttcat ttccaattga  
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420  
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480  
ggcagatctc ttgccatttg atgtgaaggg tgtttgggaa tccatggaag aaggcttgaa  
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240  
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300  
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360  
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tatttgatcc attggccact tagttctcag cctggaaagt ttccatttcc aattgatgtg  
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caagtgaaaa tgctgtggt tggaatggga tcagctcctg atttcacatg taagaaagat  
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acaaaagatg caatcattga agccatcaaa caagggtata gacactttga tactgctgnt  
240

gcttatggct cagaacaagc tcttggtgaa ggtttgaaag aagcaattga acttggcctt  
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gtcactagag aagagctttt tggtacttct aaactttggg tcactgaaaa tcctcctcac  
360

cttggtattc ctgctcttca aaaatctctc aagactcttc aattggagta cttggacttg  
420

tattgatcc attggccact tagttctcag cctggaaagt tttcatttcc aattgatgtg  
480

gcagatctct tgccatttga tgtgaagggt gtttggaat ccatggaaga aggcttgaaa  
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tatttgatcc attggccact tagttctcag cctggaaagt tttcatttcc aattgatgtg  
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gcagatctct tgccatttga tgtgaagggt gtttgggaat ccatggaaga aggcttgaaa  
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acaaaagatg caatcattga agccatcaaa caaggttata gacactttga tactgctgct  
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gcttatggct cagaacaagc tcttggtgga ggtttgaaag aagcaattga acttggcctt  
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gtcactagag aagagctttt tgttacttct aaactttggg tcactgaaaa tcatcctcac  
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cttggtgttc ctgctcttca aaaatctctc aagactcttc aattggagta cttggacttg  
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tatttgatcc attggccact tagttctcag cctggaaaagt tttcatttcc aattgatgtg  
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gcagatctct tgccatttga tgtgaagggt gtttggcaat ccatggaaga aggcttgaaa  
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tgcctgtggt tggaatggga tcagctcctg atttcacatg taagaaagat acaaaagatg  
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caatcattga agccatcaaa caagggtata gacactttga tactgctgct gcttatggct  
240

cagaacaagc tcttggtgaa ggtttgaaag aagcaattga acttggcctt gtcactagag  
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aagagctttt tggtacttct aaactttggg tcaactgaaa tcactcctcac cttggtgttc  
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ctgctcttca aaaatctctc aagactcttc aattggagta cttggacttg tatttgatcc  
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attggccact tagttctcag cctggaaagt tttcatttcc aattgatgtg gcagatctct  
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tgcctgtggt tggaatggga tcagctcctg atttcacatg taagaaagat acaaagatg  
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caatcattga agccatcaaa caagggtata gacactttga tactgctgct gcttatggct  
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caatcattga agccatcaaa caagggtata gacactttga tactgctgct gcttatggct  
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cagaacaagc tcttggtgaa ggtttgaaag aagcaattga acttggcctt gtcactagag  
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aagagctttt tgttacttct aaactttggg tcaactgaaaa tcatactcac cttgttggtc  
360

ctgctcttca aaaatctctc aagactcttc aattggagta cttggacttg tatttgatcc  
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attggccact tagttctcag cccgaaaagt tttcatttcc aattgatgtg gcagatctct  
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tgccatttga tgtgaagggt gtttggggaat ccatggaaga aggcttgaaa cttggactca  
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acttctaaac tttgggtcac tgaaaatcat cctcaccttg ttgttcctgc tcttcaaaaa  
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tctctcaaga ctottcaatt ggagtacttg gacttgtatt tgatccattg gccacttagt  
420

tctcagcctg gaaagttttc atttccaatt gatgtggcag atctcttgcc atttgatgtg  
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cggctacatt gtccacgcca ccatccaaga tctcgaggat gagaacgaga caaacattt  
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ggaagcaatg gaaggagcaa agggatcatct caaatttttc gaaatggatc ttcttaacag  
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cgactctatt gcggccgccc tgaaagggtg tgccggagtt atacatcttg catgtcctaa  
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catcattggt gaagtcaaag accccgagaa gcaaattttg gaaccggcaa ttcaaggaac  
360  
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ttggacagac ottgattatt gcaaggaaaa gaagttatac taccatttg caaagacact  
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20	25	30
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35	40	45
Glu Gly Ala Lys Gly His Leu Lys Phe Phe Glu Met Asp Leu Leu Asn		
50	55	60
Ser Asp Ser Ile Ala Ala Ala Val Lys Gly Cys Ala Gly Val Ile His		
65	70	75
Leu Ala Cys Pro Asn Ile Ile Gly Glu Val Lys Asp Pro Glu Lys Gln		
85	90	95
Ile Leu Glu Pro Ala Ile Gln Gly Thr Val Asn Val Leu Lys Val Ala		
100	105	110
Lys Glu Ala Gly Val Glu Arg Val Val Ala Thr Ser Ser Ile Ser Ala		
115	120	125
Ile Ile Pro Ser Pro Asn Trp Pro Ala Asp Lys Ile Lys Gly Glu Asp		
130	135	140
Cys Trp Thr Asp Leu Asp Tyr Cys Lys Glu Lys Lys Leu Tyr Tyr Pro		
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cggctacatt gtccacgcca ccatccaaga tctcgaggat gagaacgaga caaacattt  
180

ggaagcaatg gaaggagcaa agggatcatct caaatTTTTT gaaatggatc ttcttaacag  
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480  
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cggctacatt gtccacgcca ccatccaaga tctcaggat gagaacgaga caaacattt  
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ggaagcaatg gaaggagcaa agggatcatct caaatttttc gaaatggatc ttctcaacag  
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cgactctatt gcggccgccc tgaaagggtg tgccggaggtt atacatcttg catgtcctag  
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catcattggt gaagtcaaag accccgagaa gcaaattttg gaaccggcaa ttcaaggaac  
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gatctccgcc atcataccga gtcctaattg gccagctgat aagattaagg gagaagattg  
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180  
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240  
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420  
tctctgccat cataccgagt cctaattggc cagctgataa gattaaggga gaagattggt  
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120  
  
gtacattgt ccacgccacc atccaagatc tcgaggatga gaacgagaca aaacatttgg  
180  
  
aagcaatgga aggagcaaag ggtcatctca aatttttcga aatggatctt ctcaacagcg  
240  
  
actctattgc ggccgccgtg aaagggtgtg ccggagttat acatcttgca tgtcctagca  
300  
  
tcattggtga agtcaaagac cccgagaagc aaattttgga accggcaatt caaggaacgg  
360  
  
ttaatgtgtt gaagggtggc aaggaagcag ggggtggagcg tgtggtggcg acatcgtcga  
420  
  
tctccgcat cataccgagt cctaattggc cagctgataa gattaagggg gaagattgtt  
480  
  
ggacagacct tgattattgc aaggaaaaga agttatacta cccatttgca aagacactag  
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120

acattgtcca cgccaccatc caagatctcg aggatgagaa cgagacaaaa catttggaag  
180

caatggaagg agcaaagggt catctcaaatt tttttgaaat ggatcttctt aacagcgact  
240

ctattgcggc cgccgtgaaa ggttgtgccg gagttataca tcttgcatgt cctaacaatca  
300

ttggtgaagt caaagacccc gagaagcaaa ttttggaacc ggcaattcaa ggaacggtta  
360

atgtgttgaa ggtggcaaaag gaagcagggg tggagcgtgt ggtggcgaca tcgtcgatct  
420

ccgccatcat accgagtcct aattggccag ctgataagat taagggagaa gattgttgga  
480

cagacottga ttattgcaag gaaaagaagt tatactaccc cattgcaaag acattancag  
540

aaaaagctgg ttgggaattt gctaaagaga c  
571

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120

acatcgtcca cgccaccatc caagatctcg aggatgagaa cgagacaaaa catttggaag  
180

caatggaagg ggcaaagggt tatcttaaatt ttttcgaaat ggatcttctt aacagtgact  
240

ctattgcggc cgccgtgaaa ggttggtgccg gagttataca tcttgcatgt cctaacaatca  
300

ttggtgaagt caaagacccc gagaagcaaa ttttggaacc ggcgattcaa ggaacggtta  
360

acgtgttgaa ggtggcaaag gaagcagggg tggagcgtgt ggtggcgacg tcatcgatct  
420

ccgccatcat accgagtcct aattggccag ctgataagat taaggagaa gattgttgga  
480

cggaccttga ttattgcaag gaaaagaagt tacactaccc catcgcaaag acactagcag  
540

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120

cattgtccac gccaccatcc aagatctcga ggatgagaac gagacaaaac atttggaagc  
180

aatggaagga gcaaagggtc atctcaaatt ttttgaaatg gatcttctta acagcgactc  
240

tattgcggnc gccgtgaaag gttgtgccgg agttatacat cttgcatgng ctaacatcat  
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tggggaagcn aaagnactcn ataagnan  
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120  
catttggaag caatggaagg agcaaagggt catntnaaat ttttcgaaat ggatcttntt  
180  
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232

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120  
aaagggtcac acagtaagga ctactgtaag aaaccagat gatttggaag aggttggtta  
180  
tctaactgaa ctaagtgaag acaaagagag attgaagatt ttaaaagcag atctattggt  
240  
ggaagggagt tttgatgagg cagtgagtgg tggtgatggt gtgtttcata cagcttcccc  
300  
tgttcttggt ccacatgatg acaacattca ggttactttg attgatccat gtataaaagg  
360  
aacacaaaat gtgcttaact catgcatcaa agcaaagggtg aaacgtgtgg tgtaacatc  
420  
ttcatgctct tccataagat accgtgacga tgtgcaacaa atttctcctc ttaatgaatc  
480  
tcattggagt gattctgaat actgcaaacg ctataacctg tggtatgcat atgcaaagac  
540  
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600  
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20 25 30

Arg Asn Pro Asp Asp Leu Glu Lys Val Gly Tyr Leu Thr Glu Leu Ser  
35 40 45

Glu Asp Lys Glu Arg Leu Lys Ile Leu Lys Ala Asp Leu Leu Val Glu  
50 55 60

Gly Ser Phe Asp Glu Ala Val Ser Gly Val Asp Gly Val Phe His Thr  
65 70 75 80

Ala Ser Pro Val Leu Val Pro His Asp Asp Asn Ile Gln Val Thr Leu  
85 90 95

Ile Asp Pro Cys Ile Lys Gly Thr Gln Asn Val Leu Asn Ser Cys Ile  
100 105 110

Lys Ala Lys Val Lys Arg Val Val Leu Thr Ser Ser Cys Ser Ser Ile  
115 120 125

Arg Tyr Arg Asp Asp Val Gln Gln Ile Ser Pro Leu Asn Glu Ser His  
130 135 140

Trp Ser Asp Ser Glu Tyr Cys Lys Arg Tyr Asn Leu Trp Tyr Ala Tyr  
145 150 155 160

Ala Lys Thr Leu Gly Glu Lys Glu Ala Trp Arg Ile Ala Lys Glu Ser  
165 170 175

Gly Ile Asp Leu Val Val Val Asn Pro Ser Phe Val Gly Gly  
180 185 190

<210> 149  
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180  
tctaactgaa ctaagtgaag acaaagagag attgaagatt ttaaaagcag atctattggt  
240  
ggaagggagt tttgatgagg cagtgagtgg tgttgatggt gtgtttcata cagcttcccc  
300  
tgttcttgtt ccacatgatg acaatattca ggttactttg attgatccat gtataaaagg  
360  
aacacaaaat gtgcttagct catgcatcaa agcaaagggtg aaacgtgtgg tgtaacatc  
420  
ttcatgctct tccataagat accgtgacga tgtgcaacaa atttcaccac ttaatgaatc  
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tcattggagt gattctgaat actgcaaacg ctataacctg tggatatgcat atgcaaagac  
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attaggagaa aaagaagcat ggaggattgc  
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120

gtaagaaacc cagatgattt ggagaagggt gggtatctaa ctgaactaag tgaagacaaa  
180

gagagattga agatttttaa agcagatcta ttgggtggaag ggagttttga tgaggcagtg  
240

agtgggtgtg atgggtgtgt tcatacagct tccccgttc ttgtccaca tgatgacaat  
300

attcagggtta ctttgattga tccatgtata aaaggaacac agaattgtgt taactcatgc  
360

atcaaagcaa aggtgaaacg tgtgggtgtta acatcctcat gctcttccat aagataccgt  
420

gacgatgtgc aacaaatttc tcctcttaat gaatctcatt ggagtgttc tgattactgc  
480

aaacgctata acctgtggta tgcatatgca aagactttag gagaaaaaga agcatggagg  
540

attgcaaagg aaagtgggat taatctagtt gt  
572

&lt;210&gt; 151

&lt;211&gt; 572

&lt;212&gt; DNA

<213> *Trifolium repens*

&lt;400&gt; 151

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tgaagcctt attagaaaag ggtcacacag taaggactac tgtaagaaac ccagatgatt  
120

tggagaagggt tgggtatcta actgaactta gtgaagacaa agagagattg aagattttta  
180

aagcagatct attgggtggaa gggagttttg atgaggcagt gagtgggtgt gatgggtgtgt  
240

ttcatacagc tccccgttc cttgttccac atgatgacaa cattcagggt actttgattg  
300

atccatgtat aaaaggaaca caaatgtgc ttaactcatg catcaaagca aaggtgaaac  
360

gtgtgggtgtt aacatcttca tgctcttcca taagataccg tgacgatgtg caacaaattt  
420

ctcctcttaa tgaatctcat tggagtgtt ctgaatactg caaacgctat aacctgtgggt  
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atgcatatgc aaagacttta ggagaaaaag aagcatggag gattgcaaag gaaagtggaa  
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ttgatctagt ttagttaac ccctcttttg tt  
572

&lt;210&gt; 152

&lt;211&gt; 574

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120  
aaggttggtt atctaactga acttagtgaa gacaaagaga gattgaagat tttaaagca  
180  
gatctattgg tggaaggagg ttttgatgag gcagtgagtg gtgttgatgg tgtgtttcat  
240  
acagcttccc ctgttcttgt tccacatgat gacaacattc aggttacttt gattgatcca  
300  
tgtataaaag gaacacaaaa tgtgcttaac tcatgcatca tagcaaaggt gaaacgtgtg  
360  
gtgttaacat cttcatgctc ttccataaga taccgtgacg atgtgcaaca aatttctcct  
420  
cttaatgaat ctcatggag tgattctgaa tactgcaaac gctataacct gtggtatgca  
480  
tatgcaaaga ctttaggaga aaaagaagca tggaggattg caaaggaaag tggaattgat  
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574

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120  
atctcttcac actccacccg gaagctcaat cccggatcac aattttccct gtcgatatcc  
180  
tcgactccac cgccgtcttc tccgctatca ataactgctc aggtgtcttt catgccgctt  
240  
ctccatgtac cctcgaagat ccaactgata cgcaaaaaga gcttctagaa cctgctgtac  
300  
aaggaaccct aaatgttcta gaagcatcca ggcgcgaggt accaaaccct aattggccgg  
360  
agaaaaaggc gatcgatgag gcgtcggtga cggatgttga gtactgtaaa ttgagagggg  
420  
agtggatatct ggtgtcgaaa acggaggcgg agaaggcggc ttgggatttt cgagagaaaa  
480  
atgggtggtgt tgatgtgggg gcggnctcatc cggggacttg tttgggagag ttgatacaga

540

aggagttgaa tgcgagttca gcggnntttac agaggttgat gatgggggagt gaggataactc  
600

aagagtgnta ttggnngggg ggctgnnnat gntaaagatg n  
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Leu Val Lys Thr Leu Leu Gln Lys His Tyr Lys Ile His Ala Thr Ile  
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Phe Pro Asn Ser Asn Ala Ser His Leu Phe Thr Leu His Pro Glu Ala  
35 40 45

Gln Ser Arg Ile Thr Ile Phe Pro Val Asp Ile Leu Asp Ser Thr Ala  
50 55 60

Val Phe Ser Ala Ile Asn Asn Cys Ser Gly Val Phe His Ala Ala Ser  
65 70 75 80

Pro Cys Thr Leu Glu Asp Pro Thr Asp Pro Gln Lys Glu Leu Leu Glu  
85 90 95

Pro Ala Val Gln Gly Thr Leu Asn Val Leu Glu Ala Ser Ser Ala Gln  
100 105 110

Val Pro Asn Pro Asn Trp Pro Glu Lys Lys Ala Ile Asp Glu Ala Ser  
115 120 125

Trp Thr Asp Val Glu Tyr Cys Lys Leu Arg Gly Lys Trp Tyr Leu Val  
130 135 140

Ser Lys Thr Glu Ala Glu Lys Ala Ala Trp Asp Phe Arg Glu Lys Asn  
145 150 155 160

Gly Gly Val Asp Val Gly Ala Xaa His Pro Gly Thr Cys Leu Gly Glu  
165 170 175

Leu Ile Gln Lys Glu Leu Asn Ala Ser Ser Ala Leu Gln Arg Leu Met  
180 185 190

Met Gly Ser Glu Asp Thr Gln Glu Xaa Tyr Trp Xaa Gly Gly  
195 200 205

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120  
tcctccgcgg ctacactgtt cagccaccg tccaaaatct caatgatgag aacgaaacga  
180  
agcatctaga agctctcgaa ggagcacaaa ctaatctccg tctcttcag atcgatctcc  
240  
ttaactacga cacaatcctc gctgctgtcc gcgggtgcgt cggaattttc cacctcgctt  
300  
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Val Gln Asn Leu Asn Asp Glu Asn Glu Thr Lys His Leu Glu Ala Leu
          35          40          45

Glu Gly Ala Gln Thr Asn Leu Arg Leu Phe Gln Ile Asp Leu Leu Asn
          50          55          60

Tyr Asp Thr Ile Leu Ala Ala Val Arg Gly Cys Val Gly Ile Phe His
          65          70          75          80

Leu Ala Ser Pro Cys Thr Val Asp Lys Val His Asp Pro Gln Lys Glu
          85          90          95

Leu Leu Asp Pro Ala Ile Lys Gly Thr Leu Asn Val Leu Thr Ala Ala
          100          105          110

Lys Glu Val Gly Val Lys Arg Val Val Val Thr Ser Ser Val Ser Ala
          115          120          125

Ile Thr Pro Ser Pro Asp Trp Pro Ser Asp Val Val Lys Arg Glu Asp
          130          135          140

Cys Trp Thr Asp Val Glu Tyr Cys Lys Lys Lys Glu Leu Gly Tyr Pro
          145          150          155          160

Leu Ser Lys Thr Leu Ala Glu Lys Ala Ala Trp Asp Phe Xaa Lys Glu
          165          170          175

Asn Gly Leu Asp Val Val Xaa Val Asn Pro Xaa Thr Xaa Met Gly Pro
          180          185          190

Val Phe Pro Pro Arg His Asn Ala Ser Met Leu Met Pro Trp Glu Thr
          195          200          205

Phe Leu Lys Ala Xaa Xaa Glu Thr Phe Glu Asp Tyr Phe Met Gly Leu
          210          215          220

Ala Xaa Phe Lys Asp Xaa Ala Leu Xaa His Xaa Leu Gly Tyr Glu Asn
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180  
agcatctaga agctctcgaa ggagcacaaa ctaatctcgg tctcttccag atcgatctcc  
240  
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300  
caccttgac tgtagacaaa gttcatgac ctcagaagga gcttttggat cctgcaatta  
360  
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420  
cctcgtctgt ctcggcgatt actcctagtc ctgattggcc ttctgatggt gttaaaagag  
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120

acgccaccgt ccaaaatctc aatgatgaga acgaaacgaa gcatctagaa gctctcgaag  
180

gagcacaac taatctccgt ctcttcaga tcgatctcct taactacgac acaatcctcg  
240

ctgctgtccg cggttgcgtc ggaattttcc acctcgcttc accttgcaact gtagacaaag  
300

ttcatgatcc tcagaaggag cttttggatc ctgcaattac agggactttg aatgtgctta  
360

ctgcagctaa ggaagtaggg gtgaagcgtg tggttgttac ctgctctgtc tcggcgatta  
420

ctcctagtcc tgattggcct tctgatgttg ttaaaagaga ggattgttgg actgatgttg  
480

aatattgcaa gaaaaaagag ttgggggtatc cgttgtccaa aacattggct gaaaaagctg  
540

cgtgggattt ttncaaagaa aatgggtttgg atgttgttgn ggtgaatccc ggnactgnga  
600

tgggtcctgt ttttccacca cggcataatg caagcatgct catgccttgg gaaacttttt  
660

tgaaggctgg nnctgaaaca tttgaagact attttatggg attggccnnc tttaaagatg  
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tgaaactatc nntccttacg g  
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<210> 159

<211> 582

<212> DNA

<213> Trifolium repens

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120

gttagagatt tagggaagaa gaagaaagt gaacatttat ggaaattgga aggagcaaca  
180

gaaagactag aactaatcca agctgattta atggaagaaa atagtttcga caaagcgatc  
240

atgggatgca aagggtgtctt ccacattgcc tctccagtac tcaatcatat atcagataat  
300

cctaaggcgg aaatcttgga accggcagtc caaggtagtc taaatgtgtt gcgttcttgt  
360

aagaggaacc ccgatcttgt tcgagtgggt ctagcctcat catcttcggc tgtagagta  
420

agagctgatt ttgatccaag cataccaatt gatgaatcat cttggagctc cttggaattg  
480

tgcgagaaac tcaaggcatg gtacccaatg tcaaagacaa tggcagaaaa agcagcttgg  
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582

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<211> 190

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<213> Trifolium repens

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 Val Ile Gly Thr Val Arg Asp Leu Gly Lys Lys Lys Lys Val Glu His  
                   35                                  40                                  45  
 Leu Trp Lys Leu Glu Gly Ala Thr Glu Arg Leu Glu Leu Ile Gln Ala  
                   50                                  55                                  60  
 Asp Leu Met Glu Glu Asn Ser Phe Asp Lys Ala Ile Met Gly Cys Lys  
                   65                                  70                                  75                                  80  
 Gly Val Phe His Ile Ala Ser Pro Val Leu Asn His Ile Ser Asp Asn  
                                   85                                  90                                  95  
 Pro Lys Ala Glu Ile Leu Glu Pro Ala Val Gln Gly Thr Leu Asn Val  
                   100                                  105                                  110  
 Leu Arg Ser Cys Lys Arg Asn Pro Asp Leu Val Arg Val Val Leu Ala  
                   115                                  120                                  125  
 Ser Ser Ser Ser Ala Val Arg Val Arg Ala Asp Phe Asp Pro Ser Ile  
                   130                                  135                                  140  
 Pro Ile Asp Glu Ser Ser Trp Ser Ser Leu Glu Leu Cys Glu Lys Leu  
                   145                                  150                                  155                                  160  
 Lys Ala Trp Tyr Pro Met Ser Lys Thr Met Ala Glu Lys Ala Ala Trp  
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gtttgtgtca ccggtggtgc tggttatatt ggttctcttt tagtcaaaaa gcttttggaa  
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aagggttaca ccgttcatgc tactcttaga aacttgaagg acgaatccaa agtagatttt  
240

ttgagaggct ttccacatgc agatactaga cttatgttat ttgaagctga tatatacaaa  
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tcagatgaat tttggcccgcc aattcaaggc tgtgagtttg tttttcacct tgctactcct  
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tttcaacatc aaactgattc tcagtttaag agcatagagg aagctgcaat agcaggggta  
420

aaaagcatag ctgaaaattg cataaaatca ggaacagtga gaaaattgat atacactgga  
480

actgtaattg cttcttcttc tctgaaagat gatggaagtg gctacaaaga cttcattgat  
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572

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<211> 156



&lt;212&gt; PRT

&lt;213&gt; Trifolium repens

&lt;400&gt; 162

Met Glu Arg Ser Cys Lys Val Cys Val Thr Gly Gly Ala Gly Tyr Ile  
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Gly Ser Leu Leu Val Lys Lys Leu Leu Glu Lys Gly Tyr Thr Val His  
20 25 30

Ala Thr Leu Arg Asn Leu Lys Asp Glu Ser Lys Val Asp Phe Leu Arg  
35 40 45

Gly Phe Pro His Ala Asp Thr Arg Leu Met Leu Phe Glu Ala Asp Ile  
50 55 60

Tyr Lys Ser Asp Glu Phe Trp Pro Ala Ile Gln Gly Cys Glu Phe Val  
65 70 75 80

Phe His Leu Ala Thr Pro Phe Gln His Gln Thr Asp Ser Gln Phe Lys  
85 90 95

Ser Ile Glu Glu Ala Ala Ile Ala Gly Val Lys Ser Ile Ala Glu Asn  
100 105 110

Cys Ile Lys Ser Gly Thr Val Arg Lys Leu Ile Tyr Thr Gly Thr Val  
115 120 125

Ile Ala Ser Ser Ser Leu Lys Asp Asp Gly Ser Gly Tyr Lys Asp Phe  
130 135 140

Ile Asp Glu Thr Cys Trp Thr Pro Leu His Leu Pro  
145 150 155

&lt;210&gt; 163

&lt;211&gt; 714

&lt;212&gt; DNA

&lt;213&gt; Trifolium repens

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 180  
 cctactgccca aatactgtgt tacaggagca acaggctata ttggttcatg gcttgttgaa  
 240  
 gctcttcttc aaagagggtg cactgttcat gctactgtta gagatcctga aaaatcggtta  
 300  
 cacctcctgt cgttgtggaa aggtagtgac caattgagaa ttttccgtgc ggatttgcaa  
 360  
 gaagaaggaa gtttcgatga tgccgtaaaa ggatgtattg gtgtgttcca tgttgcagct  
 420  
 tcaatgcaat tcaatattag tgacaaagaa aacactgagg actttgttga agcaaatata  
 480  
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 540  
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 600  
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 <213> *Trifolium repens*

<400> 164

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Gln Arg Gly Cys Thr Val His Ala Thr Val Arg Asp Pro Glu Lys Ser  
 35 40 45

Leu His Leu Leu Ser Leu Trp Lys Gly Ser Asp Gln Leu Arg Ile Phe  
 50 55 60

Arg Ala Asp Leu Gln Glu Glu Gly Ser Phe Asp Asp Ala Val Lys Gly  
 65 70 75 80

Cys Ile Gly Val Phe His Val Ala Ala Ser Met Gln Phe Asn Ile Ser  
85 90 95

Asp Lys Glu Asn Thr Glu Asp Phe Val Glu Ala Asn Ile Ile Asp Pro  
100 105 110

Ala Ile Lys Gly Thr Ile Asn Leu Leu Lys Ser Cys Leu Lys Ser Asn  
115 120 125

Ser Val Lys Arg Val Val Phe Thr Ser Ser Ile Ser Thr Ile Thr Ala  
130 135 140

Lys Asp Asn Asp Gly Lys Trp Lys Pro Ile Val Asp Glu Ser Cys Gln  
145 150 155 160

Thr Lys Thr Glu Ile Leu Trp Asn Thr Gln Pro Ser Gly Trp Val Tyr  
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Ala Leu Ser Lys Leu His Ala Glu Glu Ala Ala  
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taaagaagct ataaaatgga ggaaacaaca aagatggtga aaaatagtg acaaattggt  
180

cctatagcca aatactgtgt cacaggagcc acaggctata ttggttcatt gcttgttgaa  
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120  
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180  
ctcctgtcgt tgtggaaagg tagtgaccaa ttgagaattt tccgtgcgga ttgcaagaa  
240  
gaaggaagtt tcgatgatgc cgtaaaagga tgtattggtg tgttccatgt tgcagcttca  
300  
atgcaattca atattagtga caaagaaaac actgaggact ttgttgaagc aaatataatt  
360  
gaccctgcaa tcaaaggaac cataaatctt ctcaaactcat gcttgaaatc aaattcagt  
420  
aaaagggttg ttttcacatc ttccataagt actattactg ctaaagacaa cgacggaaaa  
480  
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tcaaagaggt tgcactgttc atgctactgt tagagatcct gaaaaatcgt tacacctoct  
180

gtcgttggtg aaaggtagt accaattgag aattttccgt gcggatttgc aagaagaagg  
240

aagtttcgat gatgccgtaa aaggatgtat tgggtgtgttc catgttgacag cttcaatgca  
300

attcaatatt agtgacaaaag aaaacactga ggactttgtt gaagcaaata taattgaccc  
360

tgcaatcaaa ggaaccataa atcttctcaa atcatgcttg aaatcaaatt cagtgaaaag  
420

ggttgttttc acatcttcca taagtactat tactgctaaa gacaacgacg gaaaatggaa  
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572

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120

ggtttcatcg gatcgtggct tggttatgaga cttatagagc gtggctacac ggttcgagcc

180

accgttcgcg acccagataa catgaagaag gtgaagcatt tgggtggaact gccgggtgca  
240

aaaagcaaat tgtctctttg gaaggctgat cttgataaag aggggagttt tgatgaagca  
300

attaaaggggt gcacaggagt ttttcatggt gctacaccaa tggattttga atccaaggac  
360

cctgagaatg aagtgataaa gcctacaata aatggattaa tagacatact gaaagcatgc  
420

gaaaaggcaa aaacagttag aaaattgggt ttcacatcat cggctggaac tgtggacgtt  
480

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600

tggaagtatt cgaaagagca caacatagac tttgtctcca tcattccacc tcttggtgtt  
660

ggcccccttc ttatggcctc aatgccacct agtctaata ctgctctttc tcttatcaca  
720

ggaaatgagg cccattactc aatcataaag caagggcaat acgtccattt agatgacctt  
780

tgtcttgctc atatatttct gtatgagaat ccaaaagctc aaggagata catttgctgt  
840

tcacatgaag caaccattca tcaagttgca aaacttatta aagaaaaata cccagagtgc  
900

aatgtcccaa caaaattcaa tgatatccca gatgaattgg aaattattaa attttctaaa  
960

aagaagatca cagact  
976

&lt;210&gt; 169

&lt;211&gt; 299

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 169

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Ile Gly Ser Trp Leu Val Met Arg Leu Ile Glu Arg Gly Tyr Thr Val  
20 25 30

Arg Ala Thr Val Arg Asp Pro Asp Asn Met Lys Lys Val Lys His Leu  
35 40 45

Val Glu Leu Pro Gly Ala Lys Ser Lys Leu Ser Leu Trp Lys Ala Asp  
50 55 60

Leu Asp Lys Glu Gly Ser Phe Asp Glu Ala Ile Lys Gly Cys Thr Gly  
65 70 75 80

Val Phe His Val Ala Thr Pro Met Asp Phe Glu Ser Lys Asp Pro Glu  
85 90 95

Asn Glu Val Ile Lys Pro Thr Ile Asn Gly Leu Ile Asp Ile Leu Lys  
100 105 110

Ala Cys Glu Lys Ala Lys Thr Val Arg Lys Leu Val Phe Thr Ser Ser  
115 120 125

Ala Gly Thr Val Asp Val Thr Glu His Pro Lys Ser Ile Ile Asp Glu  
130 135 140

Thr Cys Trp Ser Asp Val Asp Phe Cys Arg Arg Val Lys Met Thr Gly  
145 150 155 160

Trp Met Tyr Phe Val Ser Lys Thr Leu Ala Glu Gln Glu Ala Trp Lys  
165 170 175

Tyr Ser Lys Glu His Asn Ile Asp Phe Val Ser Ile Ile Pro Pro Leu  
180 185 190

Val Val Gly Pro Phe Leu Met Ala Ser Met Pro Pro Ser Leu Ile Thr  
195 200 205

Ala Leu Ser Leu Ile Thr Gly Asn Glu Ala His Tyr Ser Ile Ile Lys  
210 215 220

Gln Gly Gln Tyr Val His Leu Asp Asp Leu Cys Leu Ala His Ile Phe  
225 230 235 240

Leu Tyr Glu Asn Pro Lys Ala Gln Gly Arg Tyr Ile Cys Cys Ser His  
245 250 255

Glu Ala Thr Ile His Gln Val Ala Lys Leu Ile Lys Glu Lys Tyr Pro  
260 265 270

Glu Phe Asn Val Pro Thr Lys Phe Asn Asp Ile Pro Asp Glu Leu Glu  
275 280 285

Ile Ile Lys Phe Ser Lys Lys Lys Ile Thr Asp  
290 295

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120

tgtttcatcg gatcgtggct tgttatgaga cttatagagc gtggctacac ggttcgagcc  
180

accgttcgcg acccagataa catgaagaag gtgaagcatt tgggtggaact gccgggtgca  
240

aaaagcaaat tgtctctttg gaaggctgat cttgataaag aggggagttt tgatgaagca  
300

attaaagggg gcacaggagt ttttcatggt gctacaccaa tggattttga atccaaggac  
360

cctgagaatg aagtgataaa gcctacaata aatggattaa tagacatact gaaagcatgc  
420

gaaaaggcaa aaacagttaa aaaattgggt ttacatcat cggtggaac tgtggacgtt  
480

actgaacatc caaagtctat tattgatgaa acatgctgga gtgacgttga cttttgccgt  
540

agagtcaaga tgaccggttg gatgtatttt gtttcaaaga cccctag



586

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120

gtttcatcgg gtcgtggctt gttatgagac ttatggagca tggctacact gttcgagcca  
180

ccgttcgtga cccagataac atgaagaang tgaagcattt gctggaactg ccaggtgcaa  
240

aaagcaaatt gtctcttttg aaggctgatc ttgataaaga ggggagtttt gatgaagcaa  
300

ttaaaggggtg cacaggagtt tttcatgttg ctacaccaat ggattttgag tccaaggacc  
360

ctgagaatga agtgataaag cctacaataa acggattaat agacatactg aaagcatgcg  
420

aaaaggcaaa aacagttaga aaattggttt tcacatcatc ggctggaact gtggacgtta  
480

ctgaacatcc aaagtctatt attgatgaaa catgctggag tgacgttgac ttttgccgta  
540

gagtcaaaat gaccggttgg atgtatttt  
569

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120

ttnatcggat cgtggcttgt tatgagactt atanagcgtg nctacacggc tcgagccacc  
180

gttcgcgacc cagataacat gaagaagggtg aagcatttgg tggaaactgcc ggggtgcaaaa  
240

agcaaaattgt ctcttttgaa ggctgatctt gataaagagg ggagttttga tgaagcaatt  
300

aaaggggtgca caggagtttt tnatgttgct acaccaatgg attttgaatc caaggaccct  
360

gagaatgaag tgataaagcc tacaataaat ggattaatag acatactgaa agcatgcnaa  
420

aaggcaaaaa cagttaaaaa attggttttc acatcatcgg ctgnaactgt ggacgttact  
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gaacatccaa agn  
493

<210> 173

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<212> DNA

<213> *Trifolium repens*

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120  
ttcatcggat cgtggcttgt tatgagactt atagagcgtg gctacacggg tcgagccacc  
180  
gttcgcgacc cagataacat gaagaagggt aagcatttgg tggaaactgcc gggtgcaaaa  
240  
agcaaattgt ctctttggaa ggctgatctt gataaagagg ggagttttga tgaagcaatt  
300  
aaaggggtgca caggagtttt tcatgttgct acaccaatgg attttgaatc caaggaccct  
360  
gagaatgaag tgataaagcc tacaataaat ggattaatag acatactgaa agcatgcgaa  
420  
aaggcaaaaa cagttaaaaa attgggtttc acatcatcgg ctggaactgt ggacgttact  
480  
gaacatccaa agtctattat tgatgaaaca tgctggagtg acgttgactt ttgccgtana  
540  
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580

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120

ttcatcggat cgtggcttgt tatgagactt atagagcgtg gctacacggg tcgagccacc  
180

gttcgcgacc cagataacat gaagaagggtg aagcatttgg tggaaactgcc gggtgcaaaa  
240

agcaaattgt ctctttggaa ggctgatctt gataaagagg ggagttttga tgaagcaatt  
300

aaaggggtgca caggagtttt tcatgttgct acaccaatgg attttgaatc caaggaccct  
360

gagaatgaag tgataaagcc tacaataaat ggattaatag acatactgaa agcatgcgaa  
420

aaggcaaaaa cagttagaaa attggttttc acatcatcgg ctggaactgt ggacgttact  
480

gaacatccaa agtctattat tgatgaaaca tgctggagtg acgttgactt ttgccgtaga  
540

gtcaagatga ccggttggat gtattttgtt tcaaagaccc t  
581

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gaagtgaaaa catacatggg ttccgaatca gaaatagttt gtgttaccgg agcttcaggt  
120

ttcatcggat cgtggcttgt tatgagactt atagagcgtg gctacacggt tcgagccacc  
180

gttcgcgacc cagataacat gaagaagggtg aagcatttgg tggaactgcc ggggtgcaaaa  
240

agcaaattgt ctctttggaa ggctgatctt gataaagagg ggagttttga tgaagcaatt  
300

aaaggggtgca caggagtttt tcatgttgct acaccaatgg attttgaatc caaggaccct  
360

gagaatgaag tgataaagcc tacaataaat ggattaatag acatactgaa agcatgcgaa  
420

aaggcaaaaa cagttagaaa attggttttc acatcatcgg ctggaactgt ggacgttact  
480

gaacatccaa agtctattat tgatgaaaca tgctggagtg acgttgactt ttgccgtaga  
540

gtcaagatga ccggttggat gtattttgtt tcaaagaccc tagcagagca ag  
592

<210> 176  
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60

gaagtgaaaa catacatggg ttctgaatca gaaatagttt gtgttaccgg agcatcaggt  
120

ttcatcggat cgtggccttg tatgagactt atagagcgtg gctacacggt tcgagccact  
180

gttcgcgacc cagataacat gaagaagggtg aagcatttgg tggaactgcc ggggtgcaaaa  
240

agcaaattgt ctctttggaa ggctgatctt gataaagagg ggagttttga tgaagcaatt

300

aaaggggtgca caggagtttt tcatgttgct acaccaatgg attttgaatc caaggaccct  
360

gagaatgaag tgataaagcc tacaataaat ggattaatag acatactgaa agcatgcgaa  
420

aaggcaaaaa cagttaaaaa attggttttc acatcatcgg ctggaactgt ggacgttact  
480

gaacatccaa agtctattat tgatgaaaca tgctggagtg acgttgactt ttgccgtaga  
540

gtcaagatga ccggttggat gtattttgtt tcaaagaccc tagcagagca agaagctt  
598

&lt;210&gt; 177

&lt;211&gt; 576

&lt;212&gt; DNA

<213> *Trifolium repens*

&lt;220&gt;

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&lt;222&gt; (9)..(9)

&lt;223&gt; Any nucleotide

&lt;220&gt;

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&lt;222&gt; (12)..(12)

&lt;223&gt; Any nucleotide

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&lt;221&gt; misc\_feature

&lt;222&gt; (18)..(18)

&lt;223&gt; Any nucleotide

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&lt;223&gt; Any nucleotide

&lt;400&gt; 177

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aagtgaaaat atacatgggt tccgaatcgg aaatagtttg tgttaccgga gcttcagggt  
120

tcatcgggtc gtggcttggt atgagactta tggagcgcgg ctacacgggt cgagccactg  
180

ttcgcgaccc agataacatg aagaaggatga agcatttgat ggaactgccg ggtgcaaaaa  
240

gcaaattgtc tctttggaag gctgatcttg ataaagaggg gagttttgat gaagcaatta  
300

aagggtgcac aggagttttt catgttgcta caccaatgga ttttgaatcc aaggaccctg  
360



196/390

agaacgaagt gataaagcct acaataaatg gactaataga catactgaga gcatgtgaaa  
420

aggcaaaaac aattagaaga ttggttttca catcatcagc tggaactgtg gacgtaactg  
480

aacactcaaa atcaattggt gatgaaacat gttggagtga cgttgacttt tgccgtagag  
540

tcaaaatgac cggttggatg tattttgttt caaaga  
576

<210> 178  
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aagtgaaaac atacatgggt tctgaatcag aaatagtttg tgttaccgga gcatcaggtt  
120

tcatcggatc gtggcttggt atgagactta tagagcgtgg ctacacgggt cgagccactg  
180

ttcgcgaccc agataacatg aagaaggtga agcatttggt ggaactgccg ggtgcaaaaa  
240

gcaaattgtc tctttggaag gctgatcttg ataaagaggg gagttttgat gaagcaatta  
300

aagggtgcac aggagttttt catgttgcta caccaatgga ttttgaatcc aaggaccctg  
360

agaatgaagt gataaagcct acaataaatg gattaataga catactgaaa gcatgcaaaa  
420

aggcaaaaac agttaaaaaa ttggttttca catcatcggc tggaactgtg gacgttactg  
480

aacatccaaa gtctattatt gatgaaacat gctggagtga cgttgacttt tgccgtagag  
540

tcaagatgac cggttggatg tattttgttt caaagaccct agcagag  
587

<210> 179  
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120  
  
tcatcggatc gtggcttggt atgagactta tagagcgtgg ctacacgggt cgagccactg  
180  
  
ttcgcgaccc agataacatg aagaagggtga agcatttggt ggaactgccg ggtgcaaaaa  
240  
  
gcaaattgtc tcttttgaag gctgatcttg ataaagaggg gagttttgat gaagcaatta  
300  
  
aaggggtgcac aggagttttt catgttgcta caccaatgga ttttgaatcc aaggaccctg  
360  
  
agaatgaagt gataaagcct acaataaatg gattaataga catactgaaa gcatgcgaaa  
420  
  
aggcaaaaac agttaaaaaa ttggttttca catcatcggc tggaactgtg gacgttactg  
480  
  
aacatccaaa gtctattatt gatgaaacat gctggagtga cgttgacttt tgccgtagag  
540  
  
tcaagatgac cggttggatg tatttttgttt caaagaccct agcagagcaa gaagcttggg  
600  
  
agtattctaa agagcacaac atagattttg  
630

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120  
catcgggtcg tggcttggtta tgagacttat ggagcgcggc tacacggttc gagccactgt  
180  
tcgcgaccca gataacatga agaaggtgaa gcatttgatg gaactgccgg gtgcaaaaag  
240  
caaattgtct ctttggaagg ctgatcttga taaagagggg agttttgatg aagcaattaa  
300  
agggtgcaca ggagtttttc atgttgctac accaatggat tttgaatcca aggaccctga  
360  
gaacgaagtg ataaagccta caataaatgg actaatagac atactgagag catgtgaaaa  
420  
ggcaaaaaca attagaagat tggttttcac atcatcagct ggaactgtgg acgtaactga  
480  
acactcaaaa tcaattggtg atgaaacatg ttggagtgcac gttgactttt gccgtagagt  
540  
caaaatgacc gggttgatgt attttgtttc aaagaccct  
579

<210> 181  
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120  
catcggtatcg tggcttggtta tgagacttat agagcgtggc tacacggttc gagccaccgt  
180  
tcgcgaccca gataacatga agaaggtgaa gcatttggtg gaactgccgg gtgcaaaaag  
240  
caaattgtct ctttggaagg ctgatcttga taaagagggg agttttgatg aagcaattaa  
300  
agggtgcaca ggagtttttc atgttgctac accaatggat tttgaatcca aggaccctga  
360  
gaatgaagtg ataaagccta caataaatgg attaatagac atactgaaag catgcaaaaa  
420  
ggcaaaaaca gttagaaaat tggttttcac atcatcggct ggaactgtgg acgttactga  
480  
acatccaaag tctattattg atgaaacatg ctggagtgac gttgactttt gccgtagagt  
540  
caagatgacc gggtggatgt attttgtttc aaagacccta gcagagcaag aagcttggaa  
600  
gtat  
604

<210> 182  
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<212> DNA  
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<220>  
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120

cgggtcgtgg cttgttatga gacttatgga gcgtggctac actgttcgag ccaccgttcg  
180

tgaccagat aacatgaaga aggtgaagca tttgctggaa ctgccgggtg caaaaagcaa  
240

attgtctctt tggaaggctg atcttgataa agaggggagt tttgatgaag caattaaagg  
300

gtgcacagga gtttttcatg ttgctacacc aatggatttt gaatccaagg accctgagaa  
360

tgaagtgata aagcctacaa taaacggatt aatagacata ctgaaagcat gcgaaaaggc  
420

aaaaacagtt agaaaattgg ttttcacatc atcggctgga actgtggacg ttactgaaca  
480

tccaaagtct attattgatg aaacatgctg gagtgacgtt gacttttgcc gtagagtcaa  
540

aatgaccggt tggatgtatt ttgtttcaaa gaccctagca gagcag  
586

&lt;210&gt; 183

&lt;211&gt; 586

&lt;212&gt; DNA

<213> *Trifolium repens*

&lt;220&gt;

&lt;221&gt; misc\_feature

&lt;222&gt; (2)..(2)

&lt;223&gt; Any nucleotide

&lt;220&gt;

&lt;221&gt; misc\_feature

&lt;222&gt; (11)..(11)

&lt;223&gt; Any nucleotide

&lt;400&gt; 183

gnagagaagt nacctggagt gaaaatatac atgggttctg aatcggaaat agtttgtgtt  
60

accggagctt caggtttcat cgggtcgtgg cttgttatga gacttatgga gcgtggctac  
120

actgttcgag ccaccgttcg tgaccagat aacatgaaga aggtgaagca tttgctggaa  
180

ctgccgggtg caaaaagcaa attgtctctt tggaaggctg atcttgataa agaggggagt  
240

tttgatgaag caattaaagg gtgcacagga gtttttcatg ttgctacacc aatggatttt  
300

gagccaagg accctgagaa tgaagtgata aagcctacaa taaacggatt aatagacata  
360

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ctgaaagcat gcgaaaaggc aaaaacagtt agaaaattgg ttttcacatc atcggctgga  
420

actgtggacg ttactgaaca tccaaagtct attattgatg aaacatgctg gagtgacgtt  
480

gacttttgcc gtagagtcaa aatgaccggt tggatgtatt ttgtttcaaa gaccctagca  
540

gagcaagaag cttggaagta ttcgaaagag cacaacatag actttg  
586

<210> 184  
<211> 570  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (10)..(11)  
<223> Any nucleotides

<220>  
<221> misc\_feature  
<222> (23)..(23)  
<223> Any nucleotide

<400> 184  
tactgaaagn ntgcgaaaag gcnaaaacag ttagaaaatt ggttttcaca tcacggtg  
60

gaactgtgga cgttactgaa catccaaagt ctattattga tgaaacatgc tggagtgcg  
120

ttgacttttg ccgtagagtc aaaatgaccg gttggatgta ttttgtttca aagaccctag  
180

cagagcaaga agcttggaag tattcgaaag agcacaacat agactttgtc tccatcatc  
240

cacctcttgt tgttggtccc tttcttatgg cctcaatgcc acctagtcta atcactgctc  
300

tttctcttat cacaggaaat gagggccatt actcaatcat aaagcaaggg caatacgtcc  
360

atttagatga cctttgtctt gctcatatat ttctgtatga gaatccaaaa gctcaaggga  
420

gatacatattg ctgttcacat gaagcaacca ttcataagtg tgcaaaactt attaaagaaa  
480

aataccaga gttcaatgtc ccaacaaaat tcaatgatat ccagatgaa ttggaaatta  
540

ttaaattttc taaaaagaag atcacagact  
570

<210> 185  
<211> 833  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (3)..(3)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (34)..(34)  
<223> Any nucleotide

<400> 185  
ggncataaaaa actgcactag tgtgtataag tttnttagtg aaaaaagagt gtgtaaatta  
60

acatcatggc tagtatcaaa caaattggaa acaagaaagc atgtgtgatt ggtggcactg  
120

gttttggtgc atctatgttg atcaaacagt tacttgaaaa gggttatgct gttaatacta  
180

ctgttagaga cccagatagt cctaagaaaa tatctcacct agtggcactg caaagtttgg  
240

gggaactgaa tctattttaga gcagacttaa cagttgaaga agattttgat gtccttatag  
300

caggatgtga gcttggtttt caacttgcta cacctgtgaa ctttgcttct caagatcctg  
360

agaatgacat gataaagcca gcaatcaaag gtgtgttgaa tgtgttgaaa gcaagtgcaa  
420

gagcaaaaaga agtcaaaaaga gttatcttaa catcttcggc agccgcggtg actataaatg  
480

aactcaaagg gacaggtcat gttatggatg aaaccaactg gtcagatggt gaatttctga  
540

acactgcaaa gccaccact tgggggttatc ctgcttcaaa aatgctagct gaaaaggctg  
600

catggaaatt tgctgaagaa aatgacattg atctaatac tgtgatacct agtttaacaa  
660

ctggtccttc tctcacacca gatatcccat ctagtggttg cttggcaatg tctctaataa  
720

caggcaatga tttcctcata aatgctctga aaggaatgca atttctgtcg ggttcggtat  
780

ccatcactca tgttgaggat atttgccgag ctcatatatt tctggcagag aag  
833

<210> 186  
<211> 256  
<212> PRT  
<213> Trifolium repens

<400> 186

Met Ala Ser Ile Lys Gln Ile Gly Asn Lys Lys Ala Cys Val Ile Gly  
1 5 10 15

Gly Thr Gly Phe Val Ala Ser Met Leu Ile Lys Gln Leu Leu Glu Lys  
                   20                  25                  30  
 Gly Tyr Ala Val Asn Thr Thr Val Arg Asp Pro Asp Ser Pro Lys Lys  
                   35                  40                  45  
 Ile Ser His Leu Val Ala Leu Gln Ser Leu Gly Glu Leu Asn Leu Phe  
           50                  55                  60  
 Arg Ala Asp Leu Thr Val Glu Glu Asp Phe Asp Ala Pro Ile Ala Gly  
   65                  70                  75                  80  
 Cys Glu Leu Val Phe Gln Leu Ala Thr Pro Val Asn Phe Ala Ser Gln  
                   85                  90                  95  
 Asp Pro Glu Asn Asp Met Ile Lys Pro Ala Ile Lys Gly Val Leu Asn  
                   100                  105                  110  
 Val Leu Lys Ala Ser Ala Arg Ala Lys Glu Val Lys Arg Val Ile Leu  
           115                  120                  125  
 Thr Ser Ser Ala Ala Ala Val Thr Ile Asn Glu Leu Lys Gly Thr Gly  
   130                  135                  140  
 His Val Met Asp Glu Thr Asn Trp Ser Asp Val Glu Phe Leu Asn Thr  
   145                  150                  155                  160  
 Ala Lys Pro Pro Thr Trp Gly Tyr Pro Ala Ser Lys Met Leu Ala Glu  
                   165                  170                  175  
 Lys Ala Ala Trp Lys Phe Ala Glu Glu Asn Asp Ile Asp Leu Ile Thr  
                   180                  185                  190  
 Val Ile Pro Ser Leu Thr Thr Gly Pro Ser Leu Thr Pro Asp Ile Pro  
           195                  200                  205  
 Ser Ser Val Gly Leu Ala Met Ser Leu Ile Thr Gly Asn Asp Phe Leu  
   210                  215                  220  
 Ile Asn Ala Leu Lys Gly Met Gln Phe Leu Ser Gly Ser Leu Ser Ile  
   225                  230                  235                  240  
 Thr His Val Glu Asp Ile Cys Arg Ala His Ile Phe Leu Ala Glu Lys  
                   245                  250                  255

&lt;210&gt; 187

&lt;211&gt; 576

&lt;212&gt; DNA

&lt;213&gt; Trifolium repens



<220>  
<221> misc\_feature  
<222> (3)..(3)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (34)..(34)  
<223> Any nucleotide

<400> 187  
ggncataaaa actgcactag tgtgtataag ttnatagtg aaaaagagt gtgtaaatta  
60  
acatcatggc tagtatcaaa caaattggaa acaagaaagc atgtgtgatt ggtggcactg  
120  
gttttggtgc atctatgttg atcaaacagt tacttgaaaa gggttatgct gttaatacta  
180  
ctgttagaga cccagatagt cctaagaaaa tatctcacct agtggcactg caagtttgg  
240  
gggaactgaa tctattttaga gcagacttaa cagttgaaga agattttgat gtcctatag  
300  
caggatgtga gcttggtttt caacttgcta cacctgtgaa ctttgcttct caagatcctg  
360  
agaatgacat gataaagcca gcaatcaaag gtgtgttgaa tgtgttgaaa gcaagtgcaa  
420  
gagcaaaaga agtcaaaaga gttatcttaa catcttcggc agccgcggtg actataaatg  
480  
aactcaaagg gacaggtcat gttatggatg aaaccaactg gtcagatgtt gaatttctga  
540  
acactgcaaa gccacccact tggggttatc ctgctt  
576

<210> 188  
<211> 580  
<212> DNA  
<213> Trifolium repens

<220>  
<221> misc\_feature  
<222> (3)..(3)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (5)..(5)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (34)..(35)  
<223> Any nucleotides

<220>  
<221> misc\_feature  
<222> (580)..(580)  
<223> Any nucleotide

<400> 188  
ggncntaaaa actgcactag tgtgtataag tttnttagtg aaaaaagagt gtgtaaatta  
60  
acatcatggc tagtatcaaa caaattggaa acaagaaagc atgtgtgatt ggtggcactg  
120  
gttttggtgc atctatgttg atcaaacagt tacttgaaaa gggttatgct gttaatacta  
180  
ctgttagaga cccagatagt cctaagaaaa tatctcacct agtggcactg caaagtttgg  
240  
gggaactgaa tctatttaga gcagacttaa cagttgaaga agattttgat gtcctatag  
300  
caggatgtga gcttggtttt caacttgcta cacctgtgaa ctttgcttct caagatcctg  
360  
agaatgacat gataaagcca gcaatcaaag gtgtgttgaa tgtgttgaaa gcaagtgcaa  
420  
gagcaaaaaga agtcaaaaaga gttatcttaa catcttcggc agccgcgggtg actataaatg  
480  
aactcaaagg gacaggtcac gttatggatg aaaccaactg gtcagatggt gaatttctga  
540  
acactgcaaa gccaccact tgggggttatc ctgcttcaan  
580

<210> 189  
<211> 578  
<212> DNA  
<213> Trifolium repens

<220>  
<221> misc\_feature  
<222> (14)..(14)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (29)..(30)  
<223> Any nucleotides

<400> 189  
taaaaaactgt actngtgtgt ataagtttnn tagtgaaaaa agagtgtgta aattaacatc  
60  
atggctagta tcaaacaaat tggaaacaag aaagcatgtg tgattgggtg cactgggttt  
120  
gttgcatcta tgttgatcaa acagttactt gaaaagggtt atgctgttaa tactactgtt  
180

agagacccag atagtcctaa gaaaatatct cacctagtgg cactgcaaag tttgggggaa  
240

ctgaatctat ttagagcaga cttaacagtt gaagaagatt ttgatgctcc tatagcagga  
300

tgtgagcttg tttttcaact tgctacacct gtgaactttg cttctcaaga tcctgagaat  
360

gacatgataa agccagcaat caaagggtgtg ttgaatgtgt tgaaagcaag tgcaagagca  
420

aaagaagtca aaagagttat cttaacatct tcggcagccg cggtgactat aaatgaactc  
480

aaagggacag gtcattgttat ggatgaaacc aactggctcag atgttgaatt tctgaacact  
540

gcaaagccac ccacttgggg ttatcctgct tcaaaaat  
578

<210> 190

<211> 619

<212> DNA

<213> *Trifolium repens*

<400> 190

taaaaactgc actagtgtgt ataagtttct tggtgaaaaa agagtttgta aattaacatc  
60

atggctagta tcaaacaaat tggaaacaag aaagcatgtg tgattgggtgg cactgggtttt  
120

gttgcattcta tgttgatcaa gcagttactt gaaaagggtt atgctgttaa tactaccgtt  
180

agagacccag atagccctaa gaaaatatct cacctagtgg cactgcaaag tttgggggaa  
240

ctgaatctat ttagagcaga cttaacagtt gaagaagatt ttgatgctcc tatagcagga  
300

tgtgaacttg tttttcaact tgctacacct gtgaactttg cttctcaaga tcctgagaat  
360

gacatgataa agccagcaat caaagggtgtg ttgaatgtgt tgaaagcaat tgcaagagca  
420

aaagaagtta aaagagttat cttaacatct tcggcagccg cggtgactat aaatgaactc  
480

aaagggacag gtcattgttat ggatgaaacc aactggctctg atgttgaatt tctcaacact  
540

gcaaaaccac ccacttgggg ttatcctgcc tcaaaaatgc tagctgaaaa ggctgcatgg  
600

aaatttgctg aagaaaatg  
619

<210> 191

<211> 619

<212> DNA

<213> Trifolium repens

<220>

<221> misc\_feature

<222> (12)..(12)

<223> Any nucleotide

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<222> (14)..(14)

<223> Any nucleotide

<220>

<221> misc\_feature

<222> (45)..(45)

<223> Any nucleotide

<220>

<221> misc\_feature

<222> (53)..(53)

<223> Any nucleotide

<220>

<221> misc\_feature

<222> (59)..(59)

<223> Any nucleotide

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<222> (614)..(614)

<223> Any nucleotide

<400> 191

gacctcgtgt gnantagttt cttggtgaaa aaagagtttg taaantaaca tcntggctng  
60

tatcaaaca attggaaca agaaagcatg tgtgattggt ggcactggtt ttgttgcac  
120

tatgttgatc aagcagttac ttgaaaaggg ttatgctggt aatactaccg ttagagaccc  
180

agatagccct aagaaaatat ctacactagt ggcactgcaa agtttggggg aactgaatct  
240

atttagagca gacttaacag ttgaagaaga ttttgatgct cctatagcag gatgtgaact  
300

tgtttttcaa cttgctacac ctgtgaactt tgcttctcaa gatcctgaga atgacatgat  
360

aaagccagca atcaaagggtg tgttgaatgt gttgaaagca attgcaagag caaagaagt  
420

taaaagagtt atcttaacat cttcggcagc cgcggtgact ataatgaac tcaaaggac  
480

aggatcatgt atggatgaaa ccaactgggtc tgatgttgaa tttctcaaca ctgcaaaacc  
540

accacttgg gggtatcctg cctcaaaaat gctagctgaa aaggctgcat ggaaatttgc  
600

tgaagaaaat gacnttgat  
619

<210> 192  
<211> 586  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (2)..(2)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (4)..(4)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (23)..(23)  
<223> Any nucleotide

<400> 192  
gnnacctga atctatttag agnagactta acagttgaag aagattttga tgctcctata  
60

gcaggatgtg agcttggttt tcaacttgct acacctgtga cctttgcttc tcaagatcct  
120

gagaatgaca tgataaagcc agcaatcaaa ggtgtgttga atgtgttgaa agcaagtgca  
180

agagcaaaag aagtcaaaag agttatctta acatcttcgg cagccgcggt gactataaat  
240

gaactcaaag ggacaggtca tggtatggat gaaaccaact ggtcaggtgt tgaatttctg  
300

aacactgcaa agccaccac ttgggggttat cctgcttcaa aaatgctagc tgaaaaggct  
360

gcatggaaat ttgctgaaga aaatgacatt gatctaata ctgtgatacc tagtttaaca  
420

actggtcctt ctctcacacc agatatccca tctagtgttg gcttggaat gtctctaata  
480

acaggcaatg atttctcat aaatgctctg aaaggaatgc aatttctgtc gggttcggtta  
540

tccatcactc atggtgagga tatttgccga gtcataatat ttctgg  
586

<210> 193  
<211> 567  
<212> DNA  
<213> *Trifolium repens*

<220>  
<221> misc\_feature  
<222> (55)..(55)  
<223> Any nucleotide

<400> 193  
ttaacagttg aagaagattt tgatgctcct atagcaggat gtgagcttgt ttttnaactt  
60  
  
gctacacctg tgacctttgc ttctcaagat cctgagaatg acatgataaa gccagcaatc  
120  
  
aaaggtgtgt tgaatgtgtt gaaagcaagt gcaagagcaa aagaagtcaa aagagttatc  
180  
  
ttaacatctt cggcagccgc ggtgactata aatgaactca aagggacagg tcatgttatg  
240  
  
gatgaaacca actggtcaga tgttgaattt ctgaacactg caaagccacc cacttggggg  
300  
  
tatcctgctt caaaaatgct agctgaaaag gctgcatgga aatttgctga agaaaatgac  
360  
  
attgatctaa tcaactgtgat acctagttta acaactgggc cttctctcac accagatatc  
420  
  
ccatctagtg ttggcttggc aatgtctcta ataacaggca atgatttcct cataaatgct  
480  
  
ctgaaaggaa tgcaatttct gtggggttcg ttatccatca ctcatgttga ggatatttgc  
540  
  
cgagctcata tatttctggc agagaag  
567

<210> 194  
<211> 597  
<212> DNA  
<213> *Trifolium repens*

<400> 194  
ggaaccaatt tgtcggactt ttttcccggg ttggcccgat tcgatttgca ggggtgtggtg  
60  
  
aaagagatgg atgtcttggg tccacgtttt gatagcatat ttgaaaaaat gattggtgaa  
120  
  
cgtaagaaga aggaagtgga ggggaaagaa aatgaaagta aggattttct gcagtttttg  
180  
  
ttgaatttga aggatgaggg tgattctaag actccattca caattaccca tgttaaggct  
240  
  
ctactcatgg acatggttgt ggggtggatca gacacatcct ccaacacaat tgagtttgca  
300  
  
ttggcagaaa tgatgaacaa cccagaagta atgaggaagg ttcaagagga attagaagat  
360  
  
gtagttggga aagataactt agtagaagag tctcacattc ataagctacc ctacttgc  
420

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gcagtgatga aagaaacact tcgtttacac ccagcacttc cacttttagt ccctcactgt  
480

ccaagtga aa ccaccaatgt tggaggctac acaattccaa agggatctcg tgtgtttgtg  
540

aacgtttggg ctattcatag agacccttcc atttgggaga aaccactaga atttgat  
597

&lt;210&gt; 195

&lt;211&gt; 199

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 195

Gly Thr Asn Leu Ser Asp Phe Phe Pro Gly Leu Ala Arg Phe Asp Leu  
1 5 10 15

Gln Gly Val Val Lys Glu Met Asp Val Leu Val Pro Arg Phe Asp Ser  
20 25 30

Ile Phe Glu Lys Met Ile Gly Glu Arg Lys Lys Lys Glu Val Glu Gly  
35 40 45

Lys Glu Asn Glu Ser Lys Asp Phe Leu Gln Phe Leu Leu Asn Leu Lys  
50 55 60

Asp Glu Gly Asp Ser Lys Thr Pro Phe Thr Ile Thr His Val Lys Ala  
65 70 75 80

Leu Leu Met Asp Met Val Val Gly Gly Ser Asp Thr Ser Ser Asn Thr  
85 90 95

Ile Glu Phe Ala Leu Ala Glu Met Met Asn Asn Pro Glu Val Met Arg  
100 105 110

Lys Val Gln Glu Glu Leu Glu Asp Val Val Gly Lys Asp Asn Leu Val  
115 120 125

Glu Glu Ser His Ile His Lys Leu Pro Tyr Leu His Ala Val Met Lys  
130 135 140

Glu Thr Leu Arg Leu His Pro Ala Leu Pro Leu Leu Val Pro His Cys  
145 150 155 160

Pro Ser Glu Thr Thr Asn Val Gly Gly Tyr Thr Ile Pro Lys Gly Ser  
165 170 175

Arg Val Phe Val Asn Val Trp Ala Ile His Arg Asp Pro Ser Ile Trp  
180 185 190

Glu Lys Pro Leu Glu Phe Asp

195

<210> 196  
<211> 700  
<212> DNA  
<213> *Trifolium repens*

<220>  
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<222> (2)..(2)  
<223> Any nucleotide

<220>  
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<222> (10)..(10)  
<223> Any nucleotide

<220>  
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<222> (22)..(22)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (698)..(698)  
<223> Any nucleotide

<400> 196  
gnaatccacn aatctcttga antaatacca tttctttaca agaacttaac catggtgatg  
60

atcactcaat accaaacctt cctttacaaa gaactttcta tatccttttt cattttcttg  
120

ataacccatt tcatcataag ttttctcttc aaaaaaaatc tcaaaaaact tccaccaggc  
180

ccaaaagggt ttccagttgt tgggtgcactc ccactaatgg gatccatgcc tcatgttacc  
240

ctattcaaaa tgtcacaaaa atatggtccc ataatgtacc taaaaatggg atcaaataac  
300

atggtttag catcaactcc ttcttcagcc aaagcatttc tcaaaacact tgacctaaat  
360

ttctccaata gaccgccgaa cgctggcgca actcacctag cttatgattc acaagacttg  
420

gttttcgccg actatggatc taggtggaaa ttacttagga aactaagtaa cttgcacatg  
480

ctcggcgga aagccctcga aaattggtcg aaagtctgtg agattgaaat gggtcacatg  
540

attcgtacaa tgtacgattg tagcaagaaa gacgaatccg ttgttgtggc cgaaatgttg  
600

acatatgcta tggccaatat gataggtcaa gttatattga gtcgtcgcgt gttcgagaca  
660



aaaggtagtg actcaaatga atttaaggat atgggttgntg  
700

<210> 197  
<211> 216  
<212> PRT  
<213> Trifolium repens

<220>  
<221> MISC\_FEATURE  
<222> (216)..(216)  
<223> Any amino acid

<400> 197

Met Val Met Ile Thr Gln Tyr Gln Thr Phe Leu Tyr Lys Glu Leu Ser  
1 5 10 15

Ile Ser Phe Phe Ile Phe Leu Ile Thr His Phe Ile Ile Ser Phe Leu  
20 25 30

Phe Lys Lys Asn Leu Lys Lys Leu Pro Pro Gly Pro Lys Gly Phe Pro  
35 40 45

Val Val Gly Ala Leu Pro Leu Met Gly Ser Met Pro His Val Thr Leu  
50 55 60

Phe Lys Met Ser Gln Lys Tyr Gly Pro Ile Met Tyr Leu Lys Met Gly  
65 70 75 80

Ser Asn Asn Met Val Val Ala Ser Thr Pro Ser Ser Ala Lys Ala Phe  
85 90 95

Leu Lys Thr Leu Asp Leu Asn Phe Ser Asn Arg Pro Pro Asn Ala Gly  
100 105 110

Ala Thr His Leu Ala Tyr Asp Ser Gln Asp Leu Val Phe Ala Asp Tyr  
115 120 125

Gly Ser Arg Trp Lys Leu Leu Arg Lys Leu Ser Asn Leu His Met Leu  
130 135 140

Gly Gly Lys Ala Leu Glu Asn Trp Ser Lys Val Arg Glu Ile Glu Met  
145 150 155 160

Gly His Met Ile Arg Thr Met Tyr Asp Cys Ser Lys Lys Asp Glu Ser  
165 170 175

Val Val Val Ala Glu Met Leu Thr Tyr Ala Met Ala Asn Met Ile Gly  
180 185 190

Gln Val Ile Leu Ser Arg Arg Val Phe Glu Thr Lys Gly Ser Asp Ser

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195

200

205

Asn Glu Phe Lys Asp Met Val Xaa  
210 215

<210> 198  
<211> 584  
<212> DNA  
<213> Trifolium repens

<220>  
<221> misc\_feature  
<222> (2)..(2)  
<223> Any nucleotide

<220>  
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<222> (10)..(10)  
<223> Any nucleotide

<220>  
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<222> (27)..(27)  
<223> Any nucleotide

<400> 198  
gnaatccacn aatctcttga attaatncca tttctttaca agaacttaac catggtgatg  
60  
  
atcactcaat accaaacott ctttttcaaa gaactttcta totccttttt cattttcttg  
120  
  
ataacccgtt tcatcataag ttttctcttc aaaaaaaatc tcaaaaaact tccaccaggc  
180  
  
ccaaagggtt ttccagttgt tgggtgactc ccactaatgg gatccatgcc tcatgttacc  
240  
  
ctattcaaaa tgtcacaaaa atatgggtccc ataatgtacc taaaaatggg atcaaatagc  
300  
  
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360  
  
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420  
  
gttttcgccg actatggatc taggtggaaa ttacttagga aactaagtaa cttgcacatg  
480  
  
ctcggcgga aagccctcga agattggctg aaagttcgtg agattgaaat gggtcacatg  
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120

catttcatca ttagttttct cttcaaaaaa aatctcaaaa aacttcacc aggcccaaaa  
180

ggttttcag ttgttggtgc actcccacta atgggatcca tgctcatgt taccctattc  
240

aaaatgtcac aaaaatatgg tcctataatg tacctaaaaa tgggatcaaa taacatggtt  
300

gtagcatcaa ctcttcttc agccaaagca tttctcaaaa cacttgacct aaatttctcc  
360

aataggccgg cgaacgctgg cgcaactcac ctagcttatg atccacaaga cttgggtttc  
420

gccgactatg gatctaggtg gaaattactt aggaaactaa gtaacttgca catgctcggc  
480

ggaaaagccc ttgaaaattg gtcgaaagt cgtgagattg aaatgggtca catgattcgt  
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acaatgtacg attgtagcaa gaaagacgaa tccgttggtg tggccgaaat gttgacatat  
600

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aatgtcacia aaatatgggc ccataatgta cctaaaaatg ggatcaaata acatggttgt  
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agcatcaact ctttcttcag ccaaagcatt tctcaaaaaca cttgacctaa atttctccaa  
360

tagaccgccg aacgctggcg cgactcacct agcttatgat tcacaagact tgggttttcgc  
420

cgactatgga tctaggtgga aattgcttag gaaactaagt aacttgcaca tgctcggcgg  
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aaaagccctc gaagattggt cgaaagtctg cgagattgag atgggtcaca tgattcgtac  
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ccagttgttg gtgcactccc actaatggga tccatgctc atgttacctt attcaaaatg  
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tcacaaaaat atggtcctat aatgtaccta aaaatgggat caaataacat ggttgtagca  
300

tcaactcctt cttcagccaa agcatttctc aaaacacttg acctaaattt ctccaatagg  
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tatggatcta ggtggaaatt acttaggaaa ctaagtaact tgcacatgct cggcggaaaa  
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ggtggtaaaa aggggtgggtt cattgtctct agtcatotcc aaggagaagc agtgaaggat  
480

tggagagagc tagtgacata tttttcatat ccaattaaac aaagagatta ttcaagggtg  
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ccagacaagc cagaaggatg gaaagaggta acagaaaaat acagtgaata cctaataaat  
600

ttagcttgca aactattgga agttttatca gaagcaatgg gtttagaaaa agaagctcta  
660

acaaaagcat gtgttgatat ggatcaaaaa gttgttataa attattaccc aaaatgcctt  
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gaacctgacc tcacacttgg ccttaaactg cacactgacc ctggcacaat tactcttttg  
780

cttcaagatc aagttgggtg ccttcaagct accaaagata atggtaagac gtggattaca  
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900

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1080

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1261

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 20 25 30  
 Asn Asn Phe Ser Asn Glu Ile Pro Ile Ile Ser Leu Ala Gly Ile Asp  
 35 40 45  
 Glu Val Asp Gly Arg Arg Thr Glu Ile Cys Asn Lys Ile Val Glu Ala  
 50 55 60  
 Cys Glu Asn Trp Gly Ile Phe Gln Val Val Asp His Gly Val Asp Thr  
 65 70 75 80  
 Lys Leu Val Ser Glu Met Thr Arg Phe Ala Arg Glu Phe Phe Ala Leu  
 85 90 95  
 Pro Pro Glu Glu Lys Leu Arg Phe Asp Met Ser Gly Gly Lys Lys Gly  
 100 105 110  
 Gly Phe Ile Val Ser Ser His Leu Gln Gly Glu Ala Val Lys Asp Trp  
 115 120 125  
 Arg Glu Leu Val Thr Tyr Phe Ser Tyr Pro Ile Lys Gln Arg Asp Tyr  
 130 135 140  
 Ser Arg Trp Pro Asp Lys Pro Glu Gly Trp Lys Glu Val Thr Glu Lys  
 145 150 155 160  
 Tyr Ser Glu Asn Leu Met Asn Leu Ala Cys Lys Leu Leu Glu Val Leu  
 165 170 175  
 Ser Glu Ala Met Gly Leu Glu Lys Glu Ala Leu Thr Lys Ala Cys Val  
 180 185 190  
 Asp Met Asp Gln Lys Val Val Ile Asn Tyr Tyr Pro Lys Cys Pro Glu  
 195 200 205  
 Pro Asp Leu Thr Leu Gly Leu Lys Arg His Thr Asp Pro Gly Thr Ile  
 210 215 220  
 Thr Leu Leu Leu Gln Asp Gln Val Gly Gly Leu Gln Ala Thr Lys Asp  
 225 230 235 240  
 Asn Gly Lys Thr Trp Ile Thr Val Gln Pro Val Glu Gly Ala Phe Val  
 245 250 255  
 Val Asn Leu Gly Asp His Gly His Tyr Leu Ser Asn Gly Arg Phe Lys



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260 265 270

Asn Ala Asp His Gln Ala Val Val Asn Ser Asn Tyr Ser Xaa Leu Ser  
275 280 285

Ile Ala Thr Phe Gln Asn Pro Ala Pro Asp Ala Thr Val Tyr Pro Leu  
290 295 300

Lys Ile Arg Glu Gly Glu Lys Ser Val Leu Glu Glu Pro Ile Thr Phe  
305 310 315 320

Ala Glu Met Tyr Arg Arg Lys Met Thr Lys Asp Leu Glu Ile Ala Arg  
325 330 335

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tttttcaggt tgttgatcat ggtgttgata caaaacttgt ttctgagatg acccgttttg  
360  
ctagagagtt ttttgctttg ccaccggaag agaagctccg gtttgacatg tccggtggta  
420  
aaaaggggtgg tttcattgtc tctagtcato ttcaaggaga agcagtgaag gattggagag  
480  
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acaaaagact cttgagtcaa gtttcgttag ggaagaagat gagcgtccaa aagttgccta  
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caataacttc agcaacgaga ttccaatcat ttctcttgct ggaattgatg aggttgatgg  
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tcgcagaaca gagatatgta acaagattgt tgaagcttgt gagaattggg gtatttttca  
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ggttgttgat catggtgttg atacaaagct tgtttctgag atgactcgtt ttgctagaga  
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gttttttgct ttgccgccgg aagagaagct ccggtttgac atgtccggtg gtaaaaaggg  
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tggtttcatt gtctctagtc atctccaagg agaagcagtg aaagattgga gagagctagt  
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gacatatttt tcatacccaa ttaaacaaag agattattca aggtggccag acaagccaga  
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gttatctctc acaacaaaaac actctcgagt caagtttcgt tagggaagaa gatgagcgtc  
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caaaagttgc ctacaataac ttcagcaacg agattccaat catttctctt gctggaattg  
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atgaggttga tggtcgtaga acagagatat gtaacaagat tggtgaagct tgtgagaatt  
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gggggtatatt tcagggttggt gatcatggtg ttgatacaaa acttggttct gagatgaccc  
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gttttgctag agagtttttt gctttgccac cggaagagaa gctccggttt gacatgtccg  
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gtggtaaaaa ggggtggtttc attgtctcta gtcattctca aggagaagca gtgaaggatt  
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ggagagagct agtgacatat ttttcatacc caattaaaca aagagattat tcaaggtggc

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360  
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180

cttgctggaa ttgatgaggt tgatggctgt agaacagaga tatgtaacaa gattgttgaa  
240

gcttgatgaga attgggggtat ttttcagggt gttgatcatg gtgttgatac aaaacttggt  
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tctgagatga cccgttttgc tagagagttt tttgctttgc caccggaaga gaagctccgg  
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tttgacatgt ccggtggtaa aaaggggtgt ttcattgtct ctagtcatct ccaaggagaa  
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gcagtgaagg attggagaga gctagtgaca tatttttcat acccaattaa acaaagagat  
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tattcaaggt ggccagacaa gccagaagga tggaaagagg taacagaaaa atacagtgaa  
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aacctaatga atttagcttg caaactattg gaagttttat cagaagcaat gggtttagaa  
600

aaagaagctc taacaaaagc atgtgttgat atggatcaaa aagttgttat aaattattac  
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180  
  
tgctggaatt gatgaggttg atggccgaaa aacanaaatn ttaccaaga ttgngggggc  
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300  
  
canaaanccc nttttgntaa anagtttttt gcttttcccc cggaanaaaa cctccggttt  
360  
  
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agaagatgag cgtccaaaag ttgcctacaa taacttcagc aacgagattc caatcatttc  
180  
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120



gagcgtccaa aagttgccta caataacttc agcaacgaga ttccaatcat ttctcttgct  
180

ggaattgatg aggttgatgg tcgtagaaca gagatatgta acaagattgt tgaagcttgt  
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gagaattggg gtatTTTTtca ggttggtgat catgggtgtg atacaaaact tgtttctgag  
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atgaccggtt ttgctagaga gttttttgct ttgccaccgg aagagaagct ccggtttgac  
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atgtccggtg gtaaaaaggg tggtttcatt gtctctagtc atcttcaagg agaagcagtg  
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aaggattgga gagagctagt gacatatTTT tcatacccaa ttaaacaag agattattca  
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ttgctagaga gttttttgct ttgccgccgg aagagaagct ccggtttgac atgtccggtg

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gtaaaaaggg tggtttcatt gtctctagtc atctccaagg agaagcagtg aaagattgga  
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gggagctagt gacatatattt tcatacccaa ttaaacaag agattattca aggtggccag  
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acaagccaga aggatggaaa gaagtaacag aaaaatacag tgaaaaccta atgaatttag  
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240  
gaattggggg atttttcagg ttgttgatca tgggtgttgat acaaaaacttg tttctgagat  
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gaccgtttt gctagagagt tttttgcttt gccaccggaa gagaagctcc ggtttgacat  
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gctggaattg atgaggttga tggtcgtaga acagagatat gtaacaagat tgttgaagct  
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tgtgagaatt ggggtatctt tcagggttgt gatcatgggtg ttgatacaaa acttgtttct  
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gagatgacct gttttgctag agagtttttt gctttgccac cggaagagaa gctccggttt  
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gacatgtccg gtggtaaaaa ggggtggtttc attgtctcta gtcattctcca aggagaagca  
420

gtgaaggatt ggagagagct agtgacatat tttcataacc caattaaaca aagagattat  
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tcaaggtggc cagacaagcc agaaggatgg aaagaggtaa cagaaaaata cagtgaaaac  
540

ctaataaatt tagcttgcaa actattggaa gttttatcag aagcaatggg tttagaaaaa  
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ctgagatgac ccgttttgct agagagtttt ttgctttgcc accggaagag aagctccggt  
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ttgacatgtc cggtggtaaa aagggtggtt tcattgtctc tagtcatctc caaggagaag  
420

cagtgaagga ttggagagag ctagtgacat atttttcata cccaattaaa caaagagatt  
480

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gatgaccctt tttgctagag agttttttgc tttgccaccg gaagagaagc tccggtttga  
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gaaggattgg agagagctag tgacatattt ttcataccca attaaacaaa gagattattc  
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aaggtggcca gacaagccag aaggatggaa agaggtaaca gaaaaataca gtgaaaacct  
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tgtgagaatt ggggtatttt tcagggttgtt gatcatgggtg ttgatacaaa acttgtttct  
300

gagatgaccc gttttgctag agagtttttt gctttgccac cggaagagaa gctccggttt  
360

gacatgtccg gtggtaaaaa ggggtggttc attgtctcta gtcattctca aggagaagca  
420

gtgaaggatt ggagagagct agtgacatat ttttcatacc caattaaaca aagagattat  
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tcaaggtggc cagacaagcc agaaggatgg aaagaggtaa cagaaaaata cagtgaaaac  
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atgagcgtcc aaaagttgcc tacaataact tcagcaacga gattccaatc atttctcttg  
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120  
ccaaaagttg cctacaataa cttcagcaac nagattccaa tcattttctct tgctggaatt  
180  
gatgaggttg atggtcgnag aacanagata tgtaacaaga ttgttgaagc ttgtgagaat  
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120

gagcgtccaa aagttgccta caataacttc agcaacgaga ttccaatcat ttctcttgct  
180

ggaattgatg aggttgatgg tcgcagaaca gagatatgta acaagattgt tgaagcttgt  
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gagaattggg gtatTTTTtca gggtgttgat catgggtgttg atacaaagct tgTTTTctgag  
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atgactcggtt ttgctagaga gttttttgct ttgccgccgg aagagaagct ccggtttgac  
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atgtccggtg gtaaaaaggg tggtttcatt gtctctagtc atctccaagg agaagcagtg  
420

aaagattgga gagagctagt gacatatTTTt tcataccCAA ttaaacaAag agattattca  
480

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572

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120

gagcgtccaa aagttgccta caataacttc agcaacgaga ttccaatcat ttctcttgct  
180

ggaattgatg aggttgatgg tcgtagaaca gagatatgta acaagattgt tgaagcttgt  
240

gagaattggg gtatTTTTca ggTTgttgat catggTgttg atacaaaact tgtttctgag  
300

atgacccgtt ttgctagaga gttttttgct ttgccaccgg aagagaagct ccggtttgac  
360

atgtccggtg gtaaaaaggg tggtttcatt gtctctagtc atctccaagg agaagcagtg

420

aaggattgga gagagctagt gacatatttt tcatacccaa ttaaacaag agattattca  
480

agggtggccag acaagccaga aggatggaaa gaggtaacag aaaaatacag tgaaaaccta  
540

atgaatttag cttgcaaact attggaagtt ttatc  
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&lt;211&gt; 596

&lt;212&gt; DNA

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&lt;222&gt; (15)..(15)

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&lt;400&gt; 225

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120

gtccaaaagt tgcctacaat aacttcagca acgagattcc aatcatttct cttgctggaa  
180

ttgatgaggt tgatggtcgt agaacagaga tatgtaacaa gattgttgaa gcttgatgaga  
240

attgggggtat ttttcagggtt gttgatcatg gtgttgatac aaaacttggt tctgagatga  
300

cccgttttgc tagagagttt tttgctttgc caccggaaga gaagctccgg tttgacatgt  
360

ccggtggttaa aaagggtggt ttcattgtct ctagtcatct ccaaggagaa gcagtgaagg  
420

attggagaga gctagtgaca tattttttcat acccaattaa acaaagagat tattcaaggt  
480

ggccagacaa gccagaagga tggaaagagg taacagaaaa atacagtga aacctaata  
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120  
caaaagttgc ctacaataac ttcagcaacg agattccaat cattttctctt gctggaattg  
180  
atgagggtga tggtcgtaga acagagatat gtaacaagat tgttgaagct tgtgagaatt  
240  
gggggtatattt tcagggtggt gatcatgggtg ttgatacaaa acttgtttct gagatgaccc  
300  
gttttgctag agagtttttt gctttgccac cggaagagaa gctccggttt gacatgtccg  
360  
gtggtaaaaa ggggtggttc attgtctcta gtcactcca aggagaagca gtgaaggatt  
420  
ggagagagct agtgacatat ttttcatacc caattaaaca aagagattat tcaaggtggc  
480  
cagacaagcc agaaggatgg aaagaggtaa cagaaaaata cagtgaaaac ctaatgaatt  
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600  
caaaagcatg tg  
612

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120  
aaagttgcct acaataactt cagcaacgag attccaatca tttctcttgc tggaattgat  
180  
gaggttgatg gtcgtagaac agagatatgt aacaagattg ttgaagcttg tgagaattgg  
240  
ggtatTTTTT aggttggtga tcatgggtgt gatacaaaac ttgtttctga gatgacccgt  
300  
tttgctagag agttttttgc tttgccaccg gaagagaagc tccggtttga catgtccggt  
360  
ggtaaaaagg gtggtttcat tgtctctagt catctccaag gagaagcagt gaaggattgg  
420  
agagagctag tgacatattt ttcataccca attaaacaaa gagattattc aagggtggcca  
480  
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120

aaaagttgcc tacaataact tcagcaacga gattccaatc atttctcttg ctggaattga  
180

tgaggttgat ggtcgtagaa cagagatatg taacaagatt gttgaagctt gtgagaattg  
240

gggtatTTTT caggttgttg atcatgggtg tgatacaaaa cttgtttctg agatgacccg  
300

ttttgctaga gagttttttg ctttgccacc ggaagagaag ctccggtttg acatgtccgg  
360

tggtaaaaag ggtggtttca ttgtctctag tcatctccaa ggagaagcag tgaaggattg  
420

gagagagcta gtgacatatt tttcatacc aattaaaca agagattatt caagggtggc  
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agacaagcca gaaggatgga aagaggtaac agaaaaatac agtgaaaacc taatgaattt  
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559

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120

caaaagttgc ctacaataac ttcagcaacg agattccaat catttctctt gctggaattg  
180

atgaggttga tggtcgcaga acagagatat gtaacaagat tggtgaagct tgtgagaatt  
240

ggggatTTTT tcaggttgtt gatcatgggtg ttgatacaaa gcttgtttct gagatgactc  
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gttttgctag agagtttttt gctttgccgc cggaagagaa gctccggttt gacatgtccg  
360

gtggtaaaaa ggggtggttt attgtctcta gtcattctca aggagaagca gtgaaagatt  
420

ggagagagct agtgacatat tttcatacc caattaaaca aagagattat tcaagggtggc  
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cagacaagcc agaaggatgg aaagaagtaa cagaaaaata cagtgaaaac ctaatgaatt  
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taagttatct ctcncaacaa aacactctcg agtcaagttt cgttagggaa gaagatgagc  
120

gtccaaaagt tgcctacaat aacttcagca acgagattcc aatcatttct cttgctggaa

180

ttgatgaggt tgatggtcga agaacagaga tatgtaacaa gattgttgaa gcttgtgaga  
240

attgggggtat ttttcaggtt gttgatcatg gtgttgatac aaaacttggt tctgagatga  
300

cccgttttgc tagagagttt tttgctttgc caccggaaga gaagctccgg tttgacatgt  
360

ccggtggtaa aaaggggtgtt ttcattgtct ctagtcatct ccaaggagaa gcagtgaagg  
420

attggagaga gctagtgaca tatttttcat acccaattaa acaaagagat tattcaaggt  
480

ggccagacaa gccagaagga tggaaagagg taacagaaaa atacagtga aacctaatga  
540

atthagcttg caaactattg gaagttttat cagaagcaat gggtttagaa n  
591

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<400> 231

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120

tacaaaaagt tgcctacaat aacttcagca acgagattcc aatcatttct cttgctggaa  
180

ttgatgaggt tgatggtcgt agaacagaga tatgtaacaa gattgttgaa gcttgtgaga  
240

attgggggtat ttttcagggtt gttgatcatg gtgttgatac aaaacttggt tctgagatga  
300

cccgttttgc tagagagttt tttgctttgc caccggaaga gaagctccgg tttgacatgt  
360

ccggtggtaa aaaggggtggt ttcattgtct ctagtcatct tcaaggagaa gcagtgaagg  
420

attggagaga gctagtgaca tatttttcat acccaattaa acaaagagat tattcaaggt  
480

ggccagacaa gccagaagga tggaaagagg taacagaaaa atacagtga aacctaata  
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600

taacaaaag  
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120

cctacaataa cttcagcaac gagattccaa tcattttctct tgctggaatt gatgaggttg  
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atgggtcgtag aacagagata tgtaacaaga ttgttgaagc ttgtgagaat tgggggtatctt  
240

ttcagggttg tgatcatggt gttgatacaa aacttgtttc tgagatgacc cgtttttgcta  
300

gagagttttt tgctttgcca ccggaagaga agctccggtt tgacatgtcc ggtggtaaaa  
360

agggtgggtt cattgtctct agtcatctcc aaggagaagc agtgaaggat tggagagagc  
420

tagtgacata tttttcatat ccaattaaac aaagagatta ttcaaggtgg ccagacaagc  
480

cagaaggatg gaaagaggta acagaaaaat acagtgaaaa cctaataaat ttagcttgca  
540

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597

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120

agttgcctac aataacttca gcaacgagat tccaatcatt tctcttgctg gaattgatga  
180

ggttgatggc cgtagaacag agatatgtaa caagattggt gaagcttgctg agaattgggg  
240

tattttttcag gttgttgatc atgggtgttg taaaaactt gtttctgaga tgaccggttt

300

tgctagagag ttttttgctt tgccaccgga agagaagctc cggtttgaca tgtncggtgg  
360

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120

caaaagttgc ctacaataac ttcagcaacg agattccaat catttctctt gctggaattg  
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atgaggttga tggtcgtaga acagagatat gtaacaagat tgttgaagct tgtgagaatt  
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ggggtatttt tcaggttggt gatcatggtg ttgatacaaa acttgtttct gagatgacct  
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gttttgctag agagtttttt gctttgccac cggaagagaa gctccggttt gacatgtccg  
360

gtggtaaaaa ggggtggtttc attgtctcta gtcattctca aggagaagca gtgaaggatt  
420

ggagagagct agtgacatat ttttcatacc caattaaaca aagagattat tcaagggtggc  
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120

aagttgccta caataacttc agcaacgaga ttccaatcat ttctcttgct ggaattgatg  
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aggttgatgg tcgtagaaca gagatatgta acaagattgt tgaagcttgt gagaattggg  
240

gtatTTTTca ggTgttgat catggTgttg atacaaaact tgTttctgag atgaccgTt  
300

ttgctagaga gTtttttgct ttgccaccgg aagagaagct ccggtttgac atgtccggtg  
360

gtaaaaaggg tggtttcatt gtctctagtc atctccaagg agaagcagtg aaggattgga  
420

gagagctagt gacatatTTT tcatacccaa ttaaataaag agattattca aggtggccag  
480

acaagccaga aggatggaaa gaggtaacag aaaaatacag tgaaaaccta atgaatttag  
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aagcatgt  
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ctcacaacaa aaaaccgtcg agtcaagttt cgttagggaa gaagatgagc gtccaaaagt  
120

tgcctacaat aacttcagca acgagattcc aatcatttct cttgctggaa ttgatgagggt  
180

tgatgggtcgc agaacagaga tatgtaacaa gattgttgaa gcttgtgaga attgggggtat  
240

ttttcagggtt gttgatcatg gtgttgatac aaagcttggt tctgagatga ctcgttttgc  
300

tagagagttt tttgctttgc cgccggaaga gaagctccgg tttgacatgt ccggtggtaa  
360

aaagggtggt ttcattgtct ctagtcgtct ccaaggagaa gcagtgaaag attggagaga  
420

gctagtgaca tattttttcat acccaattaa acaaagagat tattcaaggt ggccagacaa  
480

gccagaagga tggaaagaag taacanaaaa atacagtga aacctaata atttancttg  
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600

nt

602

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120  
ctacaataac ttcagcaacg agattccaat cattnctctt gctggaattg atgaggttga  
180  
tggtcgcaga acagagatat gtaacaagat tggtgaagct tgtgagaatt ggggtatctt  
240  
tcaggttggt gatcatgggtg ttgatacaaa ncttgtttct gagatgaccc gttttgctag  
300  
agagtttttt gctntgccac cggaagagaa gctccggttt gacatgtccg gtggtaaaaa  
360  
gggtgggttc attgtctcta gtcactnca agganaagca ntgaaggatt ggagagagct  
420  
agtgcacat ttttcatacc caattaaaca aagagattat tcaaggtggc cagacaagcc  
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<400> 238

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120

gttgccctaca ataacttcag caacgagatt ccaatcattt ctcttgctgg aattgatgag  
180

gttgatgggtc gcagaacaga gatatgtaac aagattgttg aagcttgtga gaattggggg  
240

atTTTTcagg ttgttgatca tgggtgttgat acaaagcttg tttctgagat gactcgTTTT  
300

gctagagagt tttttgcttt gccgccggaa gagaagctcc ggtttgacat gtccgggtgg  
360

aaaaaggggtg gtttcattgt ctctagtcac ctccaaggag aagcagtga agattggagg  
420

gagctagtga catatTTTTc ataccaatt aaacaaagag attattcaag gtggccagac  
480

aagccagaag gatggaaaga agtaacagaa aaatacagtg aaaacctaag gaatttagct  
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572

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120

ttgcctacaa taacttcagc aacgagattc caatcatttc tcttgctgga attgatgagg  
180

ttgatgggtcg aagaacagag atatgtaaca agattgttga agcttggtgag aattggggta  
240

tttttcaggt tggtgatcat ggtgttgata caaaacttgt ttctgagatg acccgttttg

300

ctagagagtt ttttgctttg ccaccggaag agaagctccg gtttgacatg tccggtggta  
360

aaaagggtgg tttcattgtc tctagtcac tccaaggaga agcagtgaag gattggagag  
420

agctagtgaac atatTTTTTca tacccaatta aacaaagaga ttattcaagg tggccagaca  
480

agccagaagg atggaaagag gtaacagaaa aatacagtga aaacctaatg aatttagctt  
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573

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120

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tgctacaat aacttcagca acgagattcc aatcatttct cttgctggaa ttgatgaggt  
180

tgatgggtcgc agaacagaga tatgtaacaa gattgttgaa gcttgtgaga attgggggtat  
240

ttttcagggtt gttgatcatg gtgttgatac aaagcttggt tctgagatga ctcgttttgc  
300

tagagagttt tttgctttgc cgccggaaga gaagctccgg tttgacatgt ccggtggtaa  
360

aaagggtggt ttcattgtct ctagtcatct ccaaggagaa gcagtgaaag attggagggga  
420

gctagtgaca tattttttcat acccaattaa acaaagagat tattcaaggt ggccagacaa  
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gccagaagga tggaaagaag taacagaaaa atacagtga aacctaata atttagcttg  
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573

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266/390

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120  
tgcttacaat aacttcagca acgagattcc aatcatttct cttgctggaa ttgatgaggt  
180  
tgatggtcgc agaacagaga tatgtaacaa gattggtgaa gcttgtgaga attgggggat  
240  
ttttcaggtt gttgatcatg gtgttgatac aaagcttggt tctgagatga ctcgttttgc  
300  
tagagagttt tttgctttgc cgccggaaga gaagctccgg tttgacatgt ccggtggtaa  
360  
aaagggtggt ttcattgtct ctagtcatct ccaaggagaa gcagtgaaag attggagaga  
420  
gctagtgaca tattttttcat acccaattaa acaaagagat tattcaaggt ggccagacaa  
480  
gccagaagga tggaaagaag taacagaaaa atacagtga aacctaata atttagcttg  
540  
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584

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tccaaaagtt gcctacaata acttcagcaa cgagattcca atnattttctc ttgctggaat  
120

tgatgagggt gatggtcgca gaacagagat atgtaacaag attggtgaag cttgtgagaa  
180

ttgggggtatt tttcagggtg ctgatcatgg tgttgatata aagcttggtt ctgagatgac  
240

tcgttttgct agagagtttt ttgctttgcc gccggaagag aagctccggt ttgacatgtc  
300

cggtgggtaaa aagggtggtt tcattgtctc tnntcatctc caaggagaag cagngaaaga  
360

ttggaganag ctagtgacat atttntcata cccaattaaa caaagagatt atncaagggtg  
420

gtcanacnag ccagaaggat ggaaagaagt aacagaaaaa tacagtgaac acctaatagaa

480

tttacctngc aagctattgg aagttttata ataancnatg gnattaaga  
529

<210> 243  
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120

ggaagagaag ctccggtttg acatgtccgg tggtaaaaag ggtggtttca ttgtctctag  
180

tcatctccaa ggagaagcag tgaaggattg gagagagcta gtgacatatt tttcataccc  
240

aattaaacaa agagattatt caaggtggcc agacaagcca gaaggatgga aagaggtaac  
300

agaaaaatac agtgaaaacc taatgaattt agcttgcaaa ctattggaag ttttatcaga  
360

agcaatgggt ttagaaaaag aagctctaac aaaagcatgt gttgatatgg atcaaaaagt  
420

tgttataaat tattacccaa aatgccctga acctgacctc acacttggcc ttaaactgca  
480

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cactgaccct ggcacaatta ctcttttgct tcaagatcaa gttggtgggc ttcaagctac  
540

caaagataat ggtaagacgt ggattacagt tcaaccagtt gaaggtgctt ttgttggttaa  
600

tcttgagac catggtcact atctaagtaa tggacgggtc aaaaatgctg accatcaagc  
660

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698

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120

ttcaagctac caaagataat ggtaagacgt ggattacagt tcaaccagtt gaaggtgctt  
180

ttgttggttaa tcttgagac catggtcatt atctaagtaa tggacgggtc aaaaatgctg  
240

accaccaagc agtggatgaat tcgaactaca gccgtttatc aatagcaaca tttcaaaatc  
300

cagctcccgat tgcaactgta taccctttga agattagaga gggtgaaaaa tctgtgttgg  
360

aagaaccaat cacttttgct gaaatgtata gaaggaagat gaccaaagac cttgaaattg  
420

ctaggatgaa gaagttggct aaggaacaac aacttaggga cttggaggag aacaagacta  
480

aatatgaggc caaacctttg aatgagatct ttgcttaatt aattagtctt aatttaaata  
540

attaataaat tttagactta atttacatat aataatttt  
579

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120

tggatcaaac aaaacaacaa ccatgggtga tctagaaaca gaaccaagtt caccatttat  
180

tcaatcccca gaacacagac caaaatcctc aataatcatt gctgaaggta tccctctaata  
240

tgatctcact cctataaact acaaagatga aatcatcacc aaccacttt ccattgaaga  
300

cttagtcaaa gaaataggca aagcatgtaa agaatggggt ttctttcaag tgattaatca  
360

caaagttcct ttggataaac gtgaaaggat tgaagaatct tcaaagaagt tttttgaact  
420

tagtttggag gaaaaactta aggtgagaag agatgaagtt aatttgcttg gttattttga  
480

agctgagcat acaaaaaatg ttagggactg gaaggaaatt tatgatttta atgtgcaaca  
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601

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<400> 246

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Thr Thr Thr Met Val Asp Leu Glu Thr Glu Pro Ser Ser Pro Phe Ile  
 20 25 30

Gln Ser Pro Glu His Arg Pro Lys Ser Ser Ile Ile Ile Ala Glu Gly  
 35 40 45

Ile Pro Leu Ile Asp Leu Thr Pro Ile Asn Tyr Lys Asp Glu Ile Ile  
 50 55 60

Thr Asn Pro Leu Ser Ile Glu Asp Leu Val Lys Glu Ile Gly Lys Ala  
 65 70 75 80

Cys Lys Glu Trp Gly Phe Phe Gln Val Ile Asn His Lys Val Pro Leu  
 85 90 95

Asp Lys Arg Glu Arg Ile Glu Glu Ser Ser Lys Lys Phe Phe Glu Leu  
 100 105 110

Ser Leu Glu Glu Lys Leu Lys Val Arg Arg Asp Glu Val Asn Leu Leu  
 115 120 125

Gly Tyr Phe Glu Ala Glu His Thr Lys Asn Val Arg Asp Trp Lys Glu  
 130 135 140

Ile Tyr Asp Phe Asn Val Gln Gln Pro Thr Phe Ile Pro Pro Ser Asp  
 145 150 155 160

Asp Gln Ser Phe Gln Phe Gln Trp Glu Asn Arg Trp  
 165 170

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120

cctgtaatca atctctcccc attaatccac cacacagttc aagactcctc tgccattgaa  
180

agcttagtca aagaaatagg aaatgcttgc aaggaatggg gtttcttcca agtaacaaac  
240

catgggtgtcc ctctaaatct aagggtcaga ctcgaggaag ctaccaaagt tttctttgca  
300

cagagtttgg aggagaagag gaagcttacc gtagatgata acagtttgcc tggttatcat  
360

gatacagagc acaccaagaa tgtcagagac tggaaagaag tgtttgattt tttatccaaa  
420

gaccccactt tgattcctct gaattctgat gaacatgatg atcgagtcac tcaatggact  
480

aatccatccc ctcaatatcc tccaaacttc aaagttattt tggaagagta tattaaagag  
540

atggaaaagc taggctttaa gttgctagag cttatagctt tgagc  
585

<210> 248

<211> 187

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<213> Trifolium repens

<400> 248

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			20					25						30	

Pro	Val	Ile	Asn	Leu	Ser	Pro	Leu	Ile	His	His	Thr	Val	Gln	Asp	Ser
		35					40						45		

Ser	Ala	Ile	Glu	Ser	Leu	Val	Lys	Glu	Ile	Gly	Asn	Ala	Cys	Lys	Glu
	50						55				60				

Trp	Gly	Phe	Phe	Gln	Val	Thr	Asn	His	Gly	Val	Pro	Leu	Asn	Leu	Arg
65					70					75					80

Leu	Arg	Leu	Glu	Glu	Ala	Thr	Lys	Val	Phe	Phe	Ala	Gln	Ser	Leu	Glu
				85					90					95	

Glu	Lys	Arg	Lys	Leu	Thr	Val	Asp	Asp	Asn	Ser	Leu	Pro	Gly	Tyr	His
			100					105					110		

Asp	Thr	Glu	His	Thr	Lys	Asn	Val	Arg	Asp	Trp	Lys	Glu	Val	Phe	Asp
		115					120					125			

Phe Leu Ser Lys Asp Pro Thr Leu Ile Pro Leu Asn Ser Asp Glu His  
 130 135 140

Asp Asp Arg Val Thr Gln Trp Thr Asn Pro Ser Pro Gln Tyr Pro Pro  
 145 150 155 160

Asn Phe Lys Val Ile Leu Glu Glu Tyr Ile Lys Glu Met Glu Lys Leu  
 165 170 175

Gly Phe Lys Leu Leu Glu Leu Ile Ala Leu Ser  
 180 185

<210> 249  
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 120  
 attcaagggtg tacaaggaaa aatgaagaaa ttacataaaa gatttgatgc atttttaact  
 180  
 agcattattg aagatcacat gatttccaag agtgagaagc ataatgactt attgagtacg  
 240  
 ttgttatcac taaaagaaaa agttgatgag gatgggtgaca aacttaatga tactgagatc  
 300  
 aaagcattac tcttgaacat gttcacagct ggaacagaca catcatcaag cacaacagag  
 360  
 tgggctattg ctgaactaat aaaaaatcca aaactaatga ttcgtgttca aaatgagttg  
 420  
 gacactgttg tgggccgaga caagcttgta actgaacaag acttggccca tcttccttac  
 480  
 ttagaggctg taataaagga gacatttcgt ctccatccat caacccctct ttctctccca  
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Val Val Glu Leu Met Ala Leu Ala Gly Val Phe Asn Ile Gly Asp Phe  
20 25 30

Val Pro Ala Leu Glu Trp Leu Asp Ile Gln Gly Val Gln Gly Lys Met  
35 40 45

Lys Lys Leu His Lys Arg Phe Asp Ala Phe Leu Thr Ser Ile Ile Glu  
50 55 60

Asp His Met Ile Ser Lys Ser Glu Lys His Asn Asp Leu Leu Ser Thr  
65 70 75 80

Leu Leu Ser Leu Lys Glu Lys Val Asp Glu Asp Gly Asp Lys Leu Asn  
85 90 95

Asp Thr Glu Ile Lys Ala Leu Leu Leu Asn Met Phe Thr Ala Gly Thr  
100 105 110

Asp Thr Ser Ser Ser Thr Thr Glu Trp Ala Ile Ala Glu Leu Ile Lys  
115 120 125

Asn Pro Lys Leu Met Ile Arg Val Gln Asn Glu Leu Asp Thr Val Val  
130 135 140

Gly Arg Asp Lys Leu Val Thr Glu Gln Asp Leu Ala His Leu Pro Tyr  
145 150 155 160

Leu Glu Ala Val Ile Lys Glu Thr Phe Arg Leu His Pro Ser Thr Pro  
165 170 175

Leu Ser Leu Pro Arg Val Ala Thr Asn Ser Cys Glu Ile Leu Asp Tyr  
180 185 190

His Ile Pro Lys Gly Ala Thr Leu Leu  
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<212> DNA

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120

atattcaagg tgtacaagga aaaatgaaga aattacataa aagatttgat gcatttttaa  
180

ctagcattat tgaagatcac atgattttcca agagtgagaa gcataatgac ttattgagta



240

cgttgttatac actaaaagaa aaagttgatg aggatgggtga caaacttaat gatactgaga  
300

tcaaagcatt actcttgaac atgttcacag ctggaacaga cacatcatca agcacaacag  
360

agtgggctat tgctgaacta ataaaaaatc caaaactaat gattcgtggt caaaatgagt  
420

tggacactgt tgtgggccga gacaagcttg taactgaaca agacttggcc catcttccct  
480

acttagaggc tgtaataaag gagacatttc gtctccatcc atcaaccctt ctttctctcc  
540

cacgtgttgc aacaaatagt tgtgaaatcc tcgactatca c  
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&lt;210&gt; 252

&lt;211&gt; 603

&lt;212&gt; DNA

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&lt;223&gt; Any nucleotide

&lt;400&gt; 252

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tggcgtttagc tggagttttc aatattgggtg attttgttcc tgctttggaa tggttagata  
120

ttcaagggtgt acaaggaaaa atgaagaaat tacataaaaag atttgatgca tttttaacta  
180

gcattattga agatcacatg atttccaaga gtgagaagca taatgactta ttgagtaagt  
240

tgttatcact aaaagaaaaa gttgatgagg atggtgacaa acttaatgat actgagatca  
300

aagcattact cttgaacatg ttcacagctg gaacagacac atcatcaagc acaacagagt  
360

gggctattgc tgaactaata aaaaatccaa aactaatgat tcgtgttcaa aatgagttgg  
420

acactgttgt gggccgagac aagcttgtaa ctgaacaaga cttggcccat cttccttact  
480

tagaggctgt aataaaggag acatttcgtc tccatccatc aaccctctt tctctccac  
540

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tgg

603

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 120  
 tatntactat tttaagatat ggaagtagta gcagcagcaa tcacaaaaaa caatggcaag  
 180  
 attgattcat tttgcttgaa tcatgctaatt gctaataaca tgaaagttaa tgggtgctgat  
 240  
 cctttgaatt ggggtgtggc tgctgaggca atgaagggaa gtcacttgga tgaggtgaag  
 300  
 cgtatgggtgg aggaataccg gaaaccggtt gtccgtcttg gtggcgagac actgacgatt  
 360  
 tctcaggtgg ctgccattgc tgcacacgat ggtgcaacgg tggagctatc ggaatctgct  
 420  
 agagccggcg ttaaggcgag cagtgactgg gttatggaga gtatgaacaa aggtacagac  
 480  
 agttatggtg tcaactacagg gtccggcgct acctcgacac gccgaaccaa acaaggtggt  
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 600  
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<400> 254

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Ser Phe Cys Leu Asn His Ala Asn Ala Asn Asn Met Lys Val Asn Gly  
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Ala Asp Pro Leu Asn Trp Gly Val Ala Ala Glu Ala Met Lys Gly Ser  
 35 40 45

His Leu Asp Glu Val Lys Arg Met Val Glu Glu Tyr Arg Lys Pro Val  
50 55 60

Val Arg Leu Gly Gly Glu Thr Leu Thr Ile Ser Gln Val Ala Ala Ile  
65 70 75 80

Ala Ala His Asp Gly Ala Thr Val Glu Leu Ser Glu Ser Ala Arg Ala  
85 90 95

Gly Val Lys Ala Ser Ser Asp Trp Val Met Glu Ser Met Asn Lys Gly  
100 105 110

Thr Asp Ser Tyr Gly Val Thr Thr Gly Phe Gly Ala Thr Ser His Arg  
115 120 125

Arg Thr Lys Gln Gly Gly Ala Leu Gln Lys Glu Leu Ile Arg Phe Glu  
130 135 140

Cys Arg Asn Ile Trp Lys Trp Asn Arg Gln Ser His Thr Leu Pro  
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120

tatntactat ttttaagatat ggaagtagta gcancagcaa tcacaaaaaa caatggcaag  
180

attgattcat tttgcttgaa tcatgctaata gctaataaca tgaaagttaa tgggtgctgat  
240

cctttgaatt ggggtgtggc tgctgaggca atgaagggaa gtcacttgga tgaggtgaag  
300

cgtatggtgg aggaataaccg gaaaccgggtt gtccgtcttg gtggcgagac gctgacgatt  
360

tctcaggtgg ctgccattgc tgcacacgat ggtgcaacgg tgganctatc ggaatctgct  
420

agagccggcg ttaaggcgag cagtgactgg gttatggaga gcatgaacaa aggtacagac  
480

agttatggtg tcactacggg gttcggcgct acctcncacc gccgaaccaa acaaggtggt  
540

gctttgcana aagagctcat aagctaattt gcttgtgtca at  
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120

tatntactat nttaagatat ggaagtagta gcagcagcaa tcacaaaaaa caatggcaag  
180

attgattcat tttgcttgaa tcatgctaatt gctaataaca tgaaagttaa tgggtgctgat  
240

cctttgaatt ggggtgtggc tgctgaggca atgaaggga gtcacttgga tgaggtgaag  
300

cgtatggtgg aggaataccg gaaaccggtt gtccgtcttg gtggcgagac actaaccatt  
360

tctcaggtgg ctgccattgc tgcacacgat ggtgcaacgg tggagctatc ggaatctgct  
420

agagccggcg ttaaggcaag cagtgactgg gttatggaga gtatgaacaa aggtaccgac  
480

agctacggtg tcccaacagg gttcggcgct acctcgacc gccgaaccaa acaaggtggt

540

gctttgcaga aagagctcat aaggtnntttt gaatgctgga atatttgga atggaacntg  
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agtcnaagcc acacactacc c  
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120tttaagnnat ggaagtagta gcagcagcaa tcacaaaaaa caacggaaag attgattcat  
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240ggggtgtggc tgctgaggca atgaaaggaa gtcacttgga tgaggtgaag tgtatggtgg  
300aggagtatcg aaaaccggtt gtccgtcttg gtggcgagac actgacgatt tctcaagtgg  
360ctgccattgc tgcacacgat ggtgcgacgg tggagctatc ggaatctgct agagccggcg  
420ttaaggcgag cagtgactgg gttatggaaa gtatgaacaa aggtactgac agttatggtg  
480tcactacagg gttcggcgct acctcgcacc gccgaaccaa acaaggtggt gctttgcaga  
540aagagctcat aaggtnntttt gaatgcagga atatttgga atggaacttg anacaaatcc  
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&lt;210&gt; 258

&lt;211&gt; 693

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&lt;222&gt; (14)..(14)

&lt;223&gt; Any nucleotide

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&lt;222&gt; (47)..(47)

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120

ctttctacac acccccctct caactattat taactaacat aatggaggga attaccaatg  
180

gccatgctga agcaactttt tgcgtgacca aaagtgttgg tgatccactc aactgggggtg  
240

cagcgcgga gtcgttgatg gggagtcat tggatgaggt gaagcgtatg gtggaggaat  
300

accgtaatcc attgggttaaa attggcggcg agacgcttac cattgctcag gtggctggaa  
360

ttgcttctca tgatagtggg gtgagggtgg agctgtctga gtccgccagg gccggcggtta  
420

aggcgagtag tgattgggtg atggacagca tgaacaatgg gactgatagt tatgggtgta  
480

ccaccggttt cggcgccacc tctcaccgga gaaccaagca gggtggtgcc ttgcagaagg  
540

agctaattag gtttttgaat gctggaatat ttggcaatgg tacagaatct aactgtacac  
600

taccacacac agcaaccaga gctgcaatgc ttgtgagaat caacactctt cttcaagagg  
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<210> 259

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<213> *Trifolium repens*

<400> 259

286/390

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Lys Ser Val Gly Asp Pro Leu Asn Trp Gly Ala Ala Ala Glu Ser Leu  
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Met Gly Ser His Leu Asp Glu Val Lys Arg Met Val Glu Glu Tyr Arg  
 35 40 45

Asn Pro Leu Val Lys Ile Gly Gly Glu Thr Leu Thr Ile Ala Gln Val  
 50 55 60

Ala Gly Ile Ala Ser His Asp Ser Gly Val Arg Val Glu Leu Ser Glu  
 65 70 75 80

Ser Ala Arg Ala Gly Val Lys Ala Ser Ser Asp Trp Val Met Asp Ser  
 85 90 95

Met Asn Asn Gly Thr Asp Ser Tyr Gly Val Thr Thr Gly Phe Gly Ala  
 100 105 110

Thr Ser His Arg Arg Thr Lys Gln Gly Gly Ala Leu Gln Lys Glu Leu  
 115 120 125

Ile Arg Phe Leu Asn Ala Gly Ile Phe Gly Asn Gly Thr Glu Ser Asn  
 130 135 140

Cys Thr Leu Pro His Thr Ala Thr Arg Ala Ala Met Leu Val Arg Ile  
 145 150 155 160

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120  
ttaactangc ataatggagg gaattaccaa tggccagctg ctttcaaaaa aaaaccaatg  
180  
gccatgctga aacaactttt tgcgtgacca aaagtgttg tgatccactc aactgggggtg  
240  
cagccgcgga gtcgttgacg ggtagtcatt tggatgaggt gaagcgtatg gtggaggagt  
300  
accgtaatcc gttgggttaa attggcggcg agacgcttac cattgctcag gtggctggaa  
360  
ttgcttctca tgatagtggg gtgagggtgg agctgtccga gtccgcaagg gccggcgcta  
420  
aggcgagtag tgattgggtg atggatagca tgaacaatgg gactgatagt tacgggtgta  
480  
ccaccggttt tgggtgccacc tctcaccgga gaaccaagca gggtggtgcc ttgcagaagg  
540  
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592

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120  
  
taatggaggg aattaccaat ggccatgctg aagcaacttt ttgcgtgacc aaaagtgttg  
180  
  
gtgatccact caactggggg gcagccgcgg agtcgttgat ggggagtcac ttggatgagg  
240  
  
tgaagcgtat ggtggaggaa taccgtaatc cattgggttaa aattggcggc gagacgctta  
300  
  
ccattgctca ggtggctgga attgcttctc atgatatggg tgtgaggggtg gagctgtctg  
360  
  
agtccgcgag ggccggcggt aaggcgagta gtgattgggt gatggacagc atgaacaatg  
420  
  
ggactgatag ttatgggtgt accaccgggt tcggtgccac ctctcaccgg agaaccaagc  
480  
  
aaggtggtgc cttgcagaag gagctaatta ggtttttgaa tgctggaata tttggcaatg  
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240  
  
tgaancgtat ggtggaggaa taccgtaatc cattgggttaa aattggcggc gagacgctta  
300  
  
ccattgctca ggtggctgga attgcttctc atgatagngn tgtgagggtg gagctgtctg  
360  
  
agtnccagcag ggccggcggt aangcgagta gtgattgngt gatggacagn atgaacaatg  
420  
  
ggactgatag ttatggngtn accaccggtt tcggcgccac ctctcaccgg agaaccaagc  
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540  
  
gtacagaatc taactgtaca cttcnacaca cagcaaccan agntgcattg ctttggtgca  
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atggagggaa ttactaatgg ccatgctgaa gcaacttttt gcgtgaccaa aagtgttggt  
180

gatccactca actgggggtgc agccgcggag tcgttgatgg ggagtcattt ggatgaggtg  
240

aagcgtatgg tggaggaata ccgtaatcca ttgggttaaaa ttggcggcga gacgcttacc  
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attgctcagg tggctggaat tgcttctcat gatagtgggtg tgagggtgga gctgtctgag  
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tccgccaggg cgggcgttaa ggcgagtagt gattgggtga tggacagcat gaacaatggg  
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actgatagtt atggtgttac caccggtttc ggcgccacct ctcaccggag aaccaagcag  
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ggtggtgcct tgcagaagga gctaattagg tttttgaatg ctggaatatt tggcaatggt  
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120

atggagggaa ttaccaatgg ccatgctgaa gcaacttttt gcgtgaccaa aagtgttggt  
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gatccactca actgggggtgc agccgcggag tcgttgatgg ggagtcattt ggatgaggtg  
240

aagcgtatgg tggaggaata ccgtaatcca ttggttaaaa ttggcggcga gacgcttacc  
300

attgctcagg tggctggaat tgcttctcat gatagtgggtg tgaggggtgga gctgtctgag  
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tccgccaggg ccggcgttaa ggcgagtagt gggtgggtga tggacagcat gaacaatggg  
420

actgatagtt atggtgttac cactggtttc ggcgccacct ctcaccggag aaccaagcag  
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ggtggtgcct tgcagaagga gctaattagg tttttgaatg ccggaatatt tggcaatggt  
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120  
  
tgagggaat taccaatggc catgctgaaa caactttttg cgtgaccaa agtggtggtg  
180  
  
atccactcaa ctggggtgca gccgcggagt cgttgacggg gagtcatttg gatgaggtga  
240  
  
agcgtatggt ggaggagtac cgtaatccgt tggctaaaat tggcggcgag acgcttacca  
300  
  
ttgctcaggt ggctggaatt gcttctcatg atagtgggtg gaggggtggag ctgnncgagt  
360  
  
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120

tttgctgac caaaagtgtt ggtgatccac tcaactgggg tgcagccgcg gagtcgtcga  
180

cggggagtca tttggatgag gtgaagcgta tggaggagga gtaccgtaat ccgttggtta  
240

aaattggcgg cgagacgctt accattgctc aggtggctgg aattgcttct catgatagtg  
300

gtgtgagggg ggagctgtcc gagtccgcaa gggccggcgt taaggcgagt agtgattggg  
360

tgatggatag catgaacaat gggactgata gttacgggtg taccaccggt tttggtgcca  
420

cctctcaccg gagaaccaag caggggtggtg ccttgcagaa ggagctaatt aggtttttga  
480

atgctggaat atttggcaat ggtacagaat ctaactttac actaccacac ac  
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120

cgatgtttca aggaacaagg ccattcatgg tggttaacttt caaggaacac ctattggagt  
180

ttcaatggat aacacacggt tagctcttgc ttcaattggt aaactcatgt ttgctcaatt  
240

ctctgaactt gttaatgatt tttacaacaa cgggttgcct tcgaatctta ctgctagtag  
300

gaacccgagc ttggactatg gtttcaaggg atcggaaatt gccatggctt cgtattgttc  
360

cgagttacaa tatcttgcta atcctgtcac caccatgtc caaagtgccg agcaacacaa  
420

ccaagatggt aactctttgg gtttgatttc atctagaaaa acaaatgaag ctattgagat



480

tctcaagctc atgtottcca ctttcttgat tgcattatgt caagcaatcg acttaaggca  
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cttggaggaa aatctcagga acaccgtcaa gaacacggt  
579

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&lt;212&gt; PRT

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Ile Glu Val Ile Arg Phe Ser Thr Lys Ser Ile Glu Arg Glu Ile Asn  
20 25 30

Ser Val Asn Asp Asn Pro Leu Ile Asp Val Ser Arg Asn Lys Ala Ile  
35 40 45

His Gly Gly Asn Phe Gln Gly Thr Pro Ile Gly Val Ser Met Asp Asn  
50 55 60

Thr Arg Leu Ala Leu Ala Ser Ile Gly Lys Leu Met Phe Ala Gln Phe  
65 70 75 80

Ser Glu Leu Val Asn Asp Phe Tyr Asn Asn Gly Leu Pro Ser Asn Leu  
85 90 95

Thr Ala Ser Arg Asn Pro Ser Leu Asp Tyr Gly Phe Lys Gly Ser Glu  
100 105 110

Ile Ala Met Ala Ser Tyr Cys Ser Glu Leu Gln Tyr Leu Ala Asn Pro  
115 120 125

Val Thr Thr His Val Gln Ser Ala Glu Gln His Asn Gln Asp Val Asn  
130 135 140

Ser Leu Gly Leu Ile Ser Ser Arg Lys Thr Asn Glu Ala Ile Glu Ile  
145 150 155 160

Leu Lys Leu Met Ser Ser Thr Phe Leu Ile Ala Leu Cys Gln Ala Ile  
165 170 175

Asp Leu Arg His Leu Glu Glu Asn Leu Arg Asn Thr Val Lys Asn Thr  
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120

caattggctg gaatcaatac cgagttcttt gaattacaac caaaagaagg tcttgcactt  
180

gttaatggaa ctgctgttgg ttctggttta gcttctattg ttctttttga ggctaacata  
240

ttggcggtgt tgtctgaagt tctatcgga attttcgctg aagttatgca agggaagccc  
300

gaatttactg atcatttgac acataagttg aagcaccacc ctggtcaaat tgaggctgct  
360

gctattatgg aacacatttt ggatgggagt gcttatgtta aagacgcaa gaagttgcat  
420

gagatggacc ctttacagaa gccaaagcaa gatagatatg cacttagaac ttcaccacaa  
480

tggcttggtc ctttgattga agtgattaga ttttcaacca agtcaattga gagagagatc  
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aactctgtca atgacaaccc tttgattgat gtttcgagaa acaaggcttt g  
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 35 40 45

Phe Phe Glu Leu Gln Pro Lys Glu Gly Leu Ala Leu Val Asn Gly Thr  
 50 55 60

Ala Val Gly Ser Gly Leu Ala Ser Ile Val Leu Phe Glu Ala Asn Ile  
 65 70 75 80

Leu Ala Val Leu Ser Glu Val Leu Ser Ala Ile Phe Ala Glu Val Met  
 85 90 95

Gln Gly Lys Pro Glu Phe Thr Asp His Leu Thr His Lys Leu Lys His  
 100 105 110

His Pro Gly Gln Ile Glu Ala Ala Ala Ile Met Glu His Ile Leu Asp  
 115 120 125

Gly Ser Ala Tyr Val Lys Asp Ala Lys Lys Leu His Glu Met Asp Pro  
 130 135 140

Leu Gln Lys Pro Lys Gln Asp Arg Tyr Ala Leu Arg Thr Ser Pro Gln  
 145 150 155 160

Trp Leu Gly Pro Leu Ile Glu Val Ile Arg Phe Ser Thr Lys Ser Ile  
 165 170 175

Glu Arg Glu Ile Asn Ser Val Asn Asp Asn Pro Leu Ile Asp Val Ser  
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Arg Asn Lys Ala Leu  
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120

ctattaagat atggaagtag tagcagcagc aatcacaaaa aacaacggca agattgattc  
180

attttgcttg aatcatgcta atgctaataa catgaaagtg aatgatgctg atcctttgaa  
240

ttgggggtgtg gctgctgagg caatgaaggg aagtcacttg gatgaggtga aacgtatggg  
300

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360

ggctgccatt gctgcacacg atgggtgcgat ggttgagctg tcggaatctg ctagagccgg  
420

cgттаaggca agcagtgatt gggttatgga gagtatgaac aaaggtactg acagttatgg  
480

tgtcaccaca gggttcggcg ctacctcnca ccgccgaacc aaacaaggtg gtgctttaca  
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Ala Asp Pro Leu Asn Trp Gly Val Ala Ala Glu Ala Met Lys Gly Ser  
 35 40 45

His Leu Asp Glu Val Lys Arg Met Val Glu Glu Tyr Arg Lys Pro Ile  
 50 55 60

Val Arg Leu Gly Gly Glu Thr Leu Thr Ile Ser Gln Val Ala Ala Ile  
 65 70 75 80

Ala Ala His Asp Gly Ala Met Val Glu Leu Ser Glu Ser Ala Arg Ala  
 85 90 95

Gly Val Lys Ala Ser Ser Asp Trp Val Met Glu Ser Met Asn Lys Gly  
 100 105 110

Thr Asp Ser Tyr Gly Val Thr Thr Gly Phe Gly Ala Thr Xaa His Arg  
 115 120 125

Arg Thr Lys Gln Gly Gly Ala Leu Gln Lys Gly Leu Ile Arg Phe Leu  
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Asn Ala Gly Ile Phe Xaa Asn Xaa Thr Xaa  
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accaatggcc atgctgaaac aacttttagc gtgacaaaa gtgnngnga tccactcaac  
180

tggcngcag ccgcggagtc gtcgacggg agtcatttgg atgaggtgaa gcgtatggng  
240

gaggagtacc gtaatccgnt ggtaaaaatt ggcggcgaga cgcttaccat tgctnnggta  
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 35 40 45

Asn Pro Xaa Val Lys Ile Gly Gly Glu Thr Leu Thr Ile Ala Xaa Val  
 50 55 60

Xaa Gly Ile Ala Ser His Asp Ser Gly Val Arg Val Glu Leu Ser Glu  
 65 70 75 80

Phe Ala Arg Ala Gly Val Lys Ala Ser Ser Asp Xaa Val Met Asp Ser  
 85 90 95

Met Asn Asn Gly Thr Asp Ser Tyr Gly Val Thr Thr Xaa Phe Gly Ala  
 100 105 110

Thr Cys His Arg Arg Thr Lys Pro Xaa Trp Cys Leu Ala Glu Gly Ala  
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cggggagtgca tttggatgag gtgaagcgta tggnggagga gtaccgtaat ccgntgggta  
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aaattggcgg cgagacgctt accattgctn nggtanctgg aattgcttct catgatagtg  
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gagtgagggg ggagctgtcc gagttcgcaa gggccggcgt taaggcgagt agtgattgng  
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tgatggatag catgaacaat gggactgata gttacgggtg taccaccgcn tttgggtgcca  
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cctgtcaccg gagaaccaag ccanggtggg gccttgacaga aggagctaaa ttnggtggtt  
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atcaagagtc ttcttgaaaa tggatactct gttaatacca ctattagagc tgatccagaa  
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cgtaagaggg atgtaagctt cctaacaaat ctacccggcg catccgaaag gctacatttc  
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ttcaacgccg atctagacga ccagagaggt ttcaacgaag caattgaagg ttgtgtcggg  
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atattccaca ccgcttcacc aatcgatttc gccgtgagtg agccagaaga aatagtgaca  
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aaaagaacag tggatggagc attaggaatt taaaagcat gtgtgaattc aaagacagtg  
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aagagattta ttacacttc aagnggttct gctgtttcat tcaatggaaa aaacaaagat  
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600

aatgggattg atgttggttac tttgattcctt cctttttattg ttggagggttt tgtttggtccc  
660

aagcttctctg attctgttga gaaagctcctt gtttttggtac taggcaaaaa ggaacaaatt  
720

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840

gaaatgtcac agcttctctc agccaaatat ccagaatata aaatactata agtagatgag  
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 Asp Leu Asp Asp Pro Glu Ser Phe Asn Glu Ala Ile Glu Gly Cys Val  
 65 70 75 80  
 Gly Ile Phe His Thr Ala Ser Pro Ile Asp Phe Ala Val Ser Glu Pro  
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 Glu Glu Ile Val Thr Lys Arg Thr Val Asp Gly Ala Leu Gly Ile Leu  
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 Lys Ala Cys Val Asn Ser Lys Thr Val Lys Arg Phe Ile Tyr Thr Ser  
 115 120 125  
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 Glu Ser Asp Trp Ser Asp Val Asp Leu Leu Arg Ser Val Lys Pro Phe  
 145 150 155 160  
 Gly Trp Ser Tyr Xaa Val Phe Lys Thr Leu Ala Glu Lys Ala Val Leu  
 165 170 175  
 Glu Phe Gly Xaa Gln Asn Gly Ile Asp Val Val Thr Leu Ile Leu Pro  
 180 185 190  
 Phe Ile Val Gly Gly Phe Val Cys Pro Lys Leu Pro Asp Ser Val Glu  
 195 200 205  
 Lys Ala Leu Val Leu Val Leu Gly Lys Lys Glu Gln Ile Gly Ile Ile  
 210 215 220  
 Ser Phe His Met Val His Val Asp Asp Val Ala Arg Ala His Ile Tyr  
 225 230 235 240  
 Leu Leu Glu Asn Pro Val Pro Gly Gly Arg Tyr Asn Cys Ser Pro Phe  
 245 250 255  
 Phe Val Ser Ile Glu Glu Met Ser Gln Leu Leu Ser Ala Lys Tyr Pro  
 260 265 270  
 Glu Tyr Gln Ile Leu Ser Val Asp Glu Leu Lys Glu Ile Lys Gly Ala  
 275 280 285

Arg Leu Pro Asp Leu Asn Ser Lys Lys Leu Val Asp Ala Gly Phe Glu  
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300  
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360  
aaaagaacag tggatggagc attaggaatt ttaaaagcat gtgtgaattc aaagacagtg  
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attctgttga gaaagctcct gttttggtac taggcaaaaa ggaacaaatt ggtattataa  
180

gtttccacat ggtacatgtg gatgatgtgg ctagagcaca tatctatcta cttgagaatc  
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ctgttccagg aggtagatat aattgttcac cattctttgt atctattgaa gaaatgtcac  
300

agcttctttc agccaaatat ccagaatatc aaatactatc tgtagatgag ttgaaggaaa  
360

ttaaaggggc aagggttgcca gatttgaact cgaagaagct cgtggacgct ggttttgagt  
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ttaagtatag tgtcgatgat atgttcgatg atgcgattca atgctgcaag gaaaaaggct  
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atctctaagc atgtgtttga aaattccatg aagttgagaa aacaatactg tgcttaaaat  
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agaaagctct tgttttggtg ctaggcaaaa aggaacaaat tgggtattata agtttccaca  
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240

gaggtagata taattgttca ccattctttg tatctattga agaaatgtca cagcttcttt  
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cagccaaata tccagaatat caaatactat ctgtagatga gttgaaggaa attaaagggg

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caaggttgcc agatttgaac tcgaagaagc tcgtggacgc tggttttgag ttttaagtata  
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gtgtcgaatga tatgttcgat gatgcgattc aatgctgcaa ggaaaaaggc tatctctaag  
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catgtgtttg aaaattccat gaagttgaga aaacaatact gtgcctaaaa tcaatgatgg  
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aggcaaaaag gaacaaattg gtattataag tttccacatg gtacatgtag atgatgtggc  
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tagagcacat atctatctac ttgagaatcc tgttccagga ggtagatata attgttcacc  
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attctttgta tctattgaag aaatgtcaca gcttctctca gccaaatata cagaatatca  
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aatactatca gtagatgagt tgaaggaaat taaagggtgca agattgccag atttgaactc  
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gaagaagctc gtggacgctg gttttgagtt taagtatagt gtcgatgata tgtttgatga  
420

tgcgattcaa tgctgcaagg aaaaaggcta tctctaagca tgtatttgaa aattccatga  
480

agttgagaaa acaataatgt gcctaaaatac aatgatggct aatgagatgt acaagtttat  
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aggcaaaaag gaacaaattg gtattataag tttccacatg gtacatgtag atgatgtggc  
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tagagcacat atctatctac ttgagaatcc tgttccagga ggtagatata attgttcacc  
240

attctttgta tctattgaag aaatgtcaca gcttctctca gccaaatata cagaatatca  
300

aatactatca gtagatgagt tgaaggaaat taaagggtgca agattgccag atttgaactc  
360

gaagaagctc gtggacgctg gttttgagtt taagtatagt gtcgatgata tgtttgatga  
420

tgcgattcaa tgctgcaagg aaaaaggcta tctctaagca tgtatttgaa aattccatga  
480

agttgagaaa acaataatgt gcctaaaatc aatgatggct aatgagatgt acaagtttat  
540

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120

agtcagagac ccaggcaatc agaagaagggt agcacacctc tggaacttag caggggccaa  
180

ggaaagggtg gagcttgtca aagctgacct cttggaagaa gggagcttcg atgatgctgt  
240

gatggcctgt gaggggtgtct tccacactgc atcacctatc atcaccaa atcgataccaa  
300

ggaagaaatg cttgattctg caattaacgg cactctaaac gtgctgagat cgtgcaagaa  
360

gaatcctttt ctcaaaaggg ttgttctcac gtcacatcg tcaaccgtga ggctgagggg  
420

tgaagctgaa ttcccaccca acgtgttgct ggatgaaaca tcatggagct ccgtggagtt  
480

ctgtgaaagt atccaggtat ggtatggtgt cgcgaagatc cttgctgaga aatcagcttg  
540

ggagttcgcc aaggagaaca acatcgacct agtggtgtgt cttccaacgt tcgtgattgg  
600

acctaattctc tegtctgaat taggaccac tgtttttagat gtccttggct tatttaaagg  
660

agagacagag aagttcacca tgtttkggaa ggatg  
695

<210> 287  
<211> 231  
<212> PRT  
<213> *Lolium perenne*

<400> 287

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Lys	Val	Ala	His	Leu	Trp	Asn	Leu	Ala	Gly	Ala	Lys	Glu	Arg	Leu	Glu
	50					55					60				
Leu	Val	Lys	Ala	Asp	Leu	Leu	Glu	Glu	Gly	Ser	Phe	Asp	Asp	Ala	Val
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Met	Ala	Cys	Glu	Gly	Val	Phe	His	Thr	Ala	Ser	Pro	Ile	Ile	Thr	Lys
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Ser	Asp	Thr	Lys	Glu	Glu	Met	Leu	Asp	Ser	Ala	Ile	Asn	Gly	Thr	Leu
			100					105					110		
Asn	Val	Leu	Arg	Ser	Cys	Lys	Lys	Asn	Pro	Phe	Leu	Lys	Arg	Val	Val
		115					120					125			
Leu	Thr	Ser	Ser	Ser	Ser	Thr	Val	Arg	Leu	Arg	Asp	Glu	Ala	Glu	Phe
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145						150					155				160
Cys	Glu	Ser	Ile	Gln	Val	Trp	Tyr	Gly	Val	Ala	Lys	Ile	Leu	Ala	Glu
				165					170					175	
Lys	Ser	Ala	Trp	Glu	Phe	Ala	Lys	Glu	Asn	Asn	Ile	Asp	Leu	Val	Ala
			180					185					190		
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		195					200					205			
Pro	Thr	Val	Leu	Asp	Val	Leu	Gly	Leu	Phe	Lys	Gly	Glu	Thr	Glu	Lys
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120  
caggcaatca gaagaaggta gcacacctct ggaacttagc aggggccaag gaaaggttgg  
180  
agcttgctcaa agctgacctc ttggaagaag ggagcttcga tgatgctgtg atggcctgtg  
240  
agggtgtctt ccacactgca tcacctatca tcaccaaato tgataccaag gaagaaatgc  
300  
ttgattctgc aattaacggc actctaaacg tgctgagatc gtgcaagaag aatccttttc  
360  
tcaaaaggggt tgttctcacg tcatcatcgt caaccgtgag gctgagggat gaagctgaat  
420  
tcccacccaa cgtgttgctg gatgaaacat catggagctc cgtggagttc tgtgaaagta  
480  
tccaggtatg gtatgggtgc gcgaagatcc ttgctgagaa atcagcttgg gagttcgcca  
540  
aggagaacaa catcgacctc gtggctgttc ttccaacggt cgtgattgga cctaattctc  
600  
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660  
agttcac  
667

<210> 289  
<211> 688  
<212> DNA  
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<220>  
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<400> 289  
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120  
gaccaggca atcagaagaa ggtagcacac ctctggaact tagcaggggc caaggaaagg  
180  
ttggagcttg tcaaagctga cctcttgga gaagggagct tcgatgatgc tgtgatggcc

240

tgtgaggggtg tcttccacac tgcacacac atcatcacca aatctgatac caaggaagaa  
300

atgcttgatt ctgcaattaa cggcactcta aacgtgctga gatcgtgcaa gaagaatcct  
360

tttctcaaaa gggttgttct cacgtcatca tcgtcaaccg tgagggtgag ggatgaagct  
420

gaattccac ccaacgtgtt gctggatgaa acatcatgga gctccgtgga gttctgtgaa  
480

agtatccagg tatggtatgg tgtcgcgaag atccttgctg agaaatcagc ttgggagttc  
540

gccaaaggaga acaacatcga cctagtggct gttcttccaa cgttcgtgat tggacctaat  
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ctctcgtctg aattaggacc cactgtttta gatgtccttg gcttatttaa nggagagaca  
660

gagaagttca ccatgttttg gaaggatg  
688

<210> 290  
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<222> (425)..(425)  
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120

gaagaaggta gcacacctct ggaacttagc agggggccaag gaaagggttg agcttggtcaa  
180

agctgacctc ttggaagaag ggagcttcga tgatgctgtg atggcctgtg aggggtgtctt  
240

ccacactgca tcacctatca tcaccaaata tgataccaag gaagaaatgc ttgattctgc  
300

aattaacggc actctaaaac ngctgagatc gngcaagaag aatncttttc tnaaaagggn  
360

tgntctcagc tcatcatcgc caccgcngan gctganggat gaanctgant tcccacccaa  
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cgngn  
425

<210> 291  
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tgttgcttcc tggcttgtca aaagacttct cgagtcgggt tataatgttc tagggacagt  
120

cagagaccca ggcaatcaga agaaggtagc acacctctgg agcttagcag gggccaagga  
180

aaggctggag cttgtcagag ctgacctctt ggaagaaggg agcttcgatg atgccgtgat  
240

ggcctgtgag ggtgtcttcc aactgcatc acctatcatc accaaatctg ataccaagga  
300

agaaatgctt gattctgcaa taaacggcac tctaaacgtg ctgagatcgt gcaagaagaa  
360

tccttttctc aaaagggttg ttctcacgtc atcatcgtca accgtgaggc tgagggatga  
420

agctgaattc ccacccaacg tgttgctgga tgaacatca tggagctccg tggagttctg  
480

tgaagtatc caggatatgt acggtgtcgc aaagatcctt gccgagaaat cagcctggga  
540

gttgccaag gagaacaaca tcgacctagt ggctgttctt ccaacattcg tgattggacc  
600

taatctctcg tctgaattag gaccactgt ttagatgtc cttggcttat ttaaaggaga  
660

gacagagaag ttcaccatgt ttgggaagga n  
691

<210> 292

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120  
gtcagagacc caggcaatca gaagaaggta gcacacctot ggagcttagc aggggccaag  
180  
gaaaggctgg agcttgctcag agctgacctc ttggaagaag ggagcttcga tgatgccgtg  
240  
atggcctgtg aggggtgtctt ccacactgca tcacctatca tcaccaaatac tgataccaag  
300  
gaagaaatgc ttgattctgc aataaacggc nctctaaacg tgctggnatc cgggtnaaaa  
360

aaan  
365

<210> 293  
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<400> 293  
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caccgcgcgc acttcctaaa gcttctcaag gacctcttcc cgcagtactc cttcaccgcc  
120  
aagtgcgaag acgacggcaa gcccatggcg aagccgtaca agttctccna ccagaggctc  
180  
agggacctgg gattaaaatt cactccgctg gcggaaagtt tgtacgagac cgtgacgtgc  
240  
ctgcaaaaaa atggccacct gcctctgccc gctcccatgg cgccaaagcg tgcataccta  
300  
taatactaca aagacacggc cgggatcgac aagccaagaa acagaggatt ctcccgaggt  
360  
tcaccatgga attgtgtatt tcacaaagtt tgaattctta ttttttttat tatgaagaaa  
420  
tacggaaaac caatactgta taccagaggc aagtgtatac atgtaaatag tcgtgtaa  
480  
cttgttcaag aatgaatgat aaagtatttt ttgcaaaaaa aaaa  
524

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<211> 100  
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Phe Pro Gln Tyr Ser Phe Thr Ala Lys Cys Glu Asp Asp Gly Lys Pro  
35 40 45  
Met Ala Lys Pro Tyr Lys Phe Ser Xaa Gln Arg Leu Arg Asp Leu Gly  
50 55 60  
Leu Lys Phe Thr Pro Leu Ala Glu Ser Leu Tyr Glu Thr Val Thr Cys  
65 70 75 80  
Leu Gln Lys Asn Gly His Leu Pro Leu Pro Ala Pro Met Ala Pro Lys  
85 90 95

Arg Ala Tyr Leu  
100

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120  
aagtgcgaag acgacggcaa gcccatggcg aagccgtaca agttctccaa ccagaggctc  
180  
agggacctgg gattaaaatt cactccgctg gcggaaagtt tgtacgagac cgtgacgtgc  
240  
ctgcaaaaaa atggccacct gcctctgccc gctcccgtgg cgccaaagcg tgcataccta  
300  
taatantacc aagacacggc cgggatcgac aagccaagaa acagaggatt ctcccgaggt  
360  
tcaccatgga attgtgtatt tcagaaagtt tgaattotta ttttttttat tatgaaggaa  
420  
tacggataac caatactgta taccagaggc aagtgttaaca atgtaaatag tcgtgtaaat  
480  
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524

<210> 296  
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<212> DNA  
<213> Lolium perenne

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<223> Any nucleotide

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<222> (25)..(26)

<223> Any nucleotides

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gcggaaagtt tgtacgagac cgtgacgtgc ctgcaaaaaa atggccacct gcctctgccc  
120

gctcccatgg cgccaaagcg tgcataccta taatactaca aagacacggc cgggatcgac  
180

aagccaagaa acagaggatt ctcccagggt tcaccatgga attgtgtatt tcacaaagtt  
240

tgaattctta ttttttttat tatgaagaaa tacggaaaac caatactgta taccagagggc  
300

aagtgtaca atgtaaatag tcgtgtaaat cttgttcaag aatgaatgat aaagtatttt  
360

ttgcaaaaaa aaaa  
374

<210> 297

<211> 363

<212> DNA

<213> Lolium perenne

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gtacgagacc gtgacgtgcc tgcaaaaaaa tggccacctg cctctgcccg ctcccatggc  
120

gccaaagcgt gcatacctat aatactacaa agacacggcc gggatcgaca agccaagaaa  
180

cagaggattc tcccagggtt caccatggaa ttgtgtattt cacaaagttt gaattcttat  
240

tttttttatt atgaagaaat acggaaaacc aatactgtat accagaggca agtgtaacaa  
300

tgtaaatagt cgtgtaaatc ttgttcaaga atgaatgata aagtattttt tgcaaaaaaa

360

aan  
363

<210> 298  
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120

agtgatcggg tggcacgctc caagaaagtc ccatctagcc acgttagagc ggtgggagac  
180

cgcccagacc tcgccaatgt cgaccacgag tccggcgcg gcatccgct catcgacctg  
240

aagcagctcg aaggtccagg gcgccgcagg gtcgtcgagg ccatcggctc cgcgtgcgag  
300

aacgatgggt ttttcatggt gacgaatcat ggcatcccag aggcggtcgt ggaggggatg  
360

ctgagcgtgg cgagggagtt cttccacctg ccggagtcgg agcggctcaa gtgctactcc  
420

gacgacccca agaaggcggc ccggctgtcg acgagcttca acgtgcgcac ggagaagggtg  
480

agcaactggc gcgacttcct ccggctgcat tgctaccctc ttgagagctt cgtcgaccag  
540

tggccgtcga acccgcccg cttcaggcaa gtcgtcggca cctactcgac ggaagcgaga  
600

gcgctggcgc tgaggctcct ggaggcgata tcggagagcc tagggctgga gagaggccac  
660

atggtgaagg ccatggggcg gcacgcgcag cacatggcgg tgaactacta cccgcccgtgc  
720

ccgcagccgg agctcaccta cggctctgcca gggcacacgg accccaacgc cctcaccatc  
780

ctcctcatgg atccccacgt ctccggcctc caggctcctca gggacggcgc caagtggatc  
840

gccgtccacc cacgccccaa cgccctggtc atcaacctag gcgaccagct acaggcgctg  
900

agcaacggcg cgtacaagag cgtgtggcac cgggcagtgg tgaacgcgga gcaggagcgt  
960

ctgtcggtag catctttcct gtgcccgtgc aacagcgcgg ttatctgccc cgcgccgagg  
1020

ctcgtcggcg acgggggagga ccccgctctac cggagctaca cctacgacga gtactacaag  
1080

aggttttgga gcaggaacct ggatcaggag cactgcctcg agctcttcag gagtcagcac  
1140

tgatgcttga accttgagtt actagctagc tctccttaac agtgcaaadc catggcccaa  
1200

gagggccccc attgcatggt tacttatggt gtttgaactg gtattgctta agtgcctaatt  
1260

aacattgcta cattctactn ctatcttgtc cgtttaaaaat tataagatgg cctaaccttt  
1320

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1380

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1381

<210> 299  
<211> 346  
<212> PRT  
<213> Lolium perenne

<400> 299

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Pro	Ser	Ser	His	Val	Arg	Ala	Val	Gly	Asp	Arg	Pro	Asp	Leu	Ala	Asn
			20					25					30		

Val	Asp	His	Glu	Ser	Gly	Ala	Gly	Ile	Pro	Leu	Ile	Asp	Leu	Lys	Gln
		35					40					45			

Leu	Glu	Gly	Pro	Gly	Arg	Arg	Val	Val	Glu	Ala	Ile	Gly	Ser	Ala	
	50				55					60					

Cys	Glu	Asn	Asp	Gly	Phe	Phe	Met	Val	Thr	Asn	His	Gly	Ile	Pro	Glu
65					70					75					80

Ala	Val	Val	Glu	Gly	Met	Leu	Ser	Val	Ala	Arg	Glu	Phe	Phe	His	Leu
				85					90					95	

Pro Glu Ser Glu Arg Leu Lys Cys Tyr Ser Asp Asp Pro Lys Lys Ala  
 100 105 110

Val Arg Leu Ser Thr Ser Phe Asn Val Arg Thr Glu Lys Val Ser Asn  
 115 120 125

Trp Arg Asp Phe Leu Arg Leu His Cys Tyr Pro Leu Glu Ser Phe Val  
 130 135 140

Asp Gln Trp Pro Ser Asn Pro Pro Ala Phe Arg Gln Val Val Gly Thr  
 145 150 155 160

Tyr Ser Thr Glu Ala Arg Ala Leu Ala Leu Arg Leu Leu Glu Ala Ile  
 165 170 175

Ser Glu Ser Leu Gly Leu Glu Arg Gly His Met Val Lys Ala Met Gly  
 180 185 190

Arg His Ala Gln His Met Ala Val Asn Tyr Tyr Pro Pro Cys Pro Gln  
 195 200 205

Pro Glu Leu Thr Tyr Gly Leu Pro Gly His Thr Asp Pro Asn Ala Leu  
 210 215 220

Thr Ile Leu Leu Met Asp Pro His Val Ser Gly Leu Gln Val Leu Arg  
 225 230 235 240

Asp Gly Ala Lys Trp Ile Ala Val His Pro Arg Pro Asn Ala Leu Val  
 245 250 255

Ile Asn Leu Gly Asp Gln Leu Gln Ala Leu Ser Asn Gly Ala Tyr Lys  
 260 265 270

Ser Val Trp His Arg Ala Val Val Asn Ala Glu Gln Glu Arg Leu Ser  
 275 280 285

Val Ala Ser Phe Leu Cys Pro Cys Asn Ser Ala Val Ile Cys Pro Ala  
 290 295 300

Pro Arg Leu Val Gly Asp Gly Glu Asp Pro Val Tyr Arg Ser Tyr Thr  
 305 310 315 320

Tyr Asp Glu Tyr Tyr Lys Arg Phe Trp Ser Arg Asn Leu Asp Gln Glu  
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His Cys Leu Glu Leu Phe Arg Ser Gln His  
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tcagctaacc attcctcaac tagaataagc atgggtccgg cgatgtccaa ccctctcctc  
120  
agtgatcggg tggcacgctc caagaaagtc ccatctagcc acgttagagc ggtgggagac  
180  
cgcccagacc tcgccaatgt cgaccacgag tccggcgcgg gcattccgct catcgacctg  
240  
aagcagctcg aagggtccagg gcgccgcagg gtctgcgagg ccatcggctc cgcggtgcgag  
300  
aacgatgggt ttttcatggg gacgaatcat ggcattcccag aggcggctcg ggaggggatg  
360  
ctgagcgtgg cgagggagtt cttccacctg ccggagtcgg agcggctcaa gtgctactcc  
420  
gacgacccca agaaggcggg ccggctgtcg acgagcttca acgtgcgcac ggagaagggtg  
480  
agcaactggc gcgacttcct ccggctgcat tgctaccctc ttgagagctt cgtcgaccag  
540  
tgcccgtcga acccgcccg cttcaggcaa gtctcgga cctactcgac ggaagcgaga  
600  
gcgctggcgc tgaggctcct ggaggcgata tcggagagcc tagggctgga gagaggccac  
660  
atgggtgaagg ccatggggcg gcacgcgcag cacatggcgg tgaactacta cccgccgtgc  
720  
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755

<210> 301  
<211> 780  
<212> DNA  
<213> Lolium perenne

<220>  
<221> misc\_feature  
<222> (21)..(21)  
<223> Any nucleotide

<220>  
<221> misc\_feature  
<222> (778)..(778)

<223> Any nucleotide

<400> 301

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60

agctaaccat tcttcaacta gaataagcat ggctccggcg atgtccaacc ctctcctcag  
120

tgatcgggtg gcacgctcca agaaagtccc atctagccac gttagagcgg tgggagaccg  
180

cccagacctc gccaatgtcg accacgagtc cggcgcgggc attccgctca tcgacctgaa  
240

gcagctcgaa ggtccagggc gccgcagggc cgtcgaggcc atcggctccg cgtgcgagaa  
300

cgatggggtt ttcatggtga cgaatcatgg catcccagag gcggtcgtgg aggggatgct  
360

gagcgtggcg agggagtctt tccacctgcc ggagtcggag cggctcaagt gctactccga  
420

cgaccccaag aaggcgggtc ggctgtcgac gagcttcaac gtgcgcacgg agaagggtgag  
480

caactggcgc gacttcctcc ggctgcattg ctacctctt gagagcttcg tcgaccagtg  
540

gccgtcgaac ccgccgcct tcaggcaagt cgtcggcacc tactcgacgg aagcgagagc  
600

gctggcgctg aggctcctgg aggcgatatc ggagagccta gggctggaga gaggccacat  
660

ggtgaaggcc atggggcggc acgcgcagca catggcgggtg aactactacc cgccgtgccc  
720

gcagccggag ctcacctacg gtctgccagg gcacacggac cccaatgccc tcaccatnct  
780

<210> 302

<211> 793

<212> DNA

<213> Lolium perenne

<220>

<221> misc\_feature

<222> (64)..(64)

<223> Any nucleotide

<220>

<221> misc\_feature

<222> (692)..(692)

<223> Any nucleotide

<220>

<221> misc\_feature

<222> (750)..(750)

<223> Any nucleotide

&lt;400&gt; 302

ggggaagtcc ggctgctggg gcttcggctt ctgggcgcga tctcgctggg gctggggctg  
60gagngagggt atgtggagaa ggtgctgggc gagcaggagc agcacatggc cgtgaactac  
120taccgcgggt gccccgagcc ggacctcacc tacggcctgc ccaagcacac ggaccccaac  
180gccctcacca tcctcctcat ggatccccac gtctccggcc tccaggctct cagggacggc  
240gccaagtgga tcgccgtcca cccacgcccc aacgccctgg tcatcaacct aggcgaccag  
300ctacaggcgc tgagcaacgg cgcgtacaag agcgtgtggc accgggcagt ggtgaacgcg  
360gagcaggagc gtctgtcggg ggcattcttc ctgtgcccg gcaacagcgc ggttatctgc  
420cccgccgcca ggctcgctcg cgacggggag gacccgtct accggagcta cacctacgac  
480gagtactaca agaggttttg gagcaggaac ctggatcagg agcactgcct cgagctcttc  
540aggagtcagc actgatgctt gaaccttgag ttactagcta gctctcctta acagtgcaaa  
600tccatggccc aagagggccc cgattgcatg gttacttatg ttgtttgaac tggatttgct  
660taagtgccta ataacattgc tacattctac tnctatcttg tccgtttaaa attataagat  
720ggcctaacct ttttcttaat tgtatgcatn ctgaacatat ttaagtgtgt gtgttcagac  
780agtttagtct gca  
793

&lt;210&gt; 303

&lt;211&gt; 1395

&lt;212&gt; DNA

&lt;213&gt; Lolium perenne

&lt;400&gt; 303

gaattcgatt aagcagtggg aacaacgcag agtacgcggg gactctcaga acacactgtg  
60taaccacggg agcgagtggc aagactagca gaaagtacgg acatcagcta accattcctc  
120aactagaata agcatggctc cggcgatgtc caaccctctc ctcagtgatc ggggtggcacg  
180ctccaagaaa gtcccatcta gccacgttag agcggtgagg gaccgcccag acctcgccaa  
240tgtcgaccac gagtccggcg cgggcattcc gctcatcgac ctgaagcagc tcgaaggctc  
300

agggcgccgc aggggtcgtcg aggccatcgg ctccgcgtgc gagaacgatg ggtttttcat  
360  
ggtgacgaat catggcatcc cagaggcggg cgtggagggg atgctgagcg tggcgagggg  
420  
gttcttccac ctgccggagt cggagcggct caagtgtac tccgacgacc ccaagaaggc  
480  
ggtccggctg tcgacgagct tcaacgtgcg cacggagaag gtgagcaact ggcgcgactt  
540  
cctccggctg cattgtacc ctcttgagag ctctgtcgac cagtggccgt cgaacccgcc  
600  
cgccttcagg caagtgtcg gcacctactc gacggaagcg agagcgctgg cgctgaggct  
660  
cctggaggcg atatcggaga gcctagggct ggagagaggc cacatggtga aggccatggg  
720  
gcggcacgcg cagcacatgg cggatgaacta ctacccgccg tgcccgagc cggagctcac  
780  
ctacggtctg ccagggcaca aggaccccaa tgccatcacg ctctctctgc aggacggcgt  
840  
ctccggcctg cagggtccagc gcgacggccg gtgggtggcc gtcaacccgg tgcccaacgc  
900  
cctcgatcatc aacatcggcg atcagttaca ggcgctgagc aacgaccgat acaagagcgt  
960  
gaaccacaga gtgatcgta acagcgcgag cgagaggatt tcggtgccga cgttctactg  
1020  
cccgtcgccg gacacgggtg tcgcgccggc cgacgcgctg gtggacgacg cccaccctcg  
1080  
ggcctaccag cccttcacgt accaggagta ctacgaggag ttctggaaga tgggccttca  
1140  
gtcagcaagt tgcctcgaca gggtccgacg gatcgagtga tggacaagac gtgggcccgtt  
1200  
gttatctcct gggccatgag cgttgccgca gccgatgtgt cgccatatgg tggagacgtt  
1260  
tcctccctcc ggaaaagaaa aataaaacag agtggagacc actagaaccg tcagatagca  
1320  
tccccaaaaa aaaaaaaaaa aaaaaaaaaa aaaagtactc tgcgttggtta ccaactgtta  
1380  
atcactagtg aattc  
1395

<210> 304  
<211> 348  
<212> PRT  
<213> Lolium perenne

<400> 304

Met Ala Pro Ala Met Ser Asn Pro Leu Leu Ser Asp Arg Val Ala Arg



1	5	10	15
Ser Lys Lys Val Pro Ser Ser His Val Arg Ala Val Gly Asp Arg Pro	20	25	30
Asp Leu Ala Asn Val Asp His Glu Ser Gly Ala Gly Ile Pro Leu Ile	35	40	45
Asp Leu Lys Gln Leu Glu Gly Pro Gly Arg Arg Arg Val Val Glu Ala	50	55	60
Ile Gly Ser Ala Cys Glu Asn Asp Gly Phe Phe Met Val Thr Asn His	65	70	75
Gly Ile Pro Glu Ala Val Val Glu Gly Met Leu Ser Val Ala Arg Glu	85	90	95
Phe Phe His Leu Pro Glu Ser Glu Arg Leu Lys Cys Tyr Ser Asp Asp	100	105	110
Pro Lys Lys Ala Val Arg Leu Ser Thr Ser Phe Asn Val Arg Thr Glu	115	120	125
Lys Val Ser Asn Trp Arg Asp Phe Leu Arg Leu His Cys Tyr Pro Leu	130	135	140
Glu Ser Phe Val Asp Gln Trp Pro Ser Asn Pro Pro Ala Phe Arg Gln	145	150	155
Val Val Gly Thr Tyr Ser Thr Glu Ala Arg Ala Leu Ala Leu Arg Leu	165	170	175
Leu Glu Ala Ile Ser Glu Ser Leu Gly Leu Glu Arg Gly His Met Val	180	185	190
Lys Ala Met Gly Arg His Ala Gln His Met Ala Val Asn Tyr Tyr Pro	195	200	205
Pro Cys Pro Gln Pro Glu Leu Thr Tyr Gly Leu Pro Gly His Lys Asp	210	215	220
Pro Asn Ala Ile Thr Leu Leu Leu Gln Asp Gly Val Ser Gly Leu Gln	225	230	235
Val Gln Arg Asp Gly Arg Trp Val Ala Val Asn Pro Val Pro Asn Ala	245	250	255
Leu Val Ile Asn Ile Gly Asp Gln Leu Gln Ala Leu Ser Asn Asp Arg	260	265	270

Tyr Lys Ser Val Asn His Arg Val Ile Val Asn Ser Ala Ser Glu Arg  
 275 280 285

Ile Ser Val Pro Thr Phe Tyr Cys Pro Ser Pro Asp Thr Val Val Ala  
 290 295 300

Pro Ala Asp Ala Leu Val Asp Asp Ala His Pro Arg Ala Tyr Gln Pro  
 305 310 315 320

Phe Thr Tyr Gln Glu Tyr Tyr Glu Glu Phe Trp Lys Met Gly Leu Gln  
 325 330 335

Ser Ala Ser Cys Leu Asp Arg Phe Arg Arg Ile Glu  
 340 345

<210> 305

<211> 1309

<212> DNA

<213> Trifolium repens

<400> 305

gaattcgatt aagcagtggg aacaacgcag agtacgcggg ataaaaactg cactagtgtg  
 60

tataagtttc ttggtgaaaa aagagtttgt aaattaacat catggctagt atcaaacaaa  
 120

ttggaacaa gaaagcatgt gtgattgggtg gcaactgggtt tgttgcattc atgttgatca  
 180

agcagttact tgaaaagggt tatgctgtta atactaccgt tagagacca gatagcccta  
 240

agaaaatatc tcacctagtg gcaactgcaa gtttggggga actgaatcta tttagagcag  
 300

acttaacagt tgaagaagat tttgatgctc ctatagcagg atgtgaactt gtttttcaac  
 360

ttgctacacc tgtgaacttt gcttctcaag atcctgagaa tgacatgata aagccagcaa  
 420

tcaaagggtg gttgaatgtg ttgaaagcaa ttgcaagagc aaaagaagtt aaaagagtta  
 480

tcttaacatc ttcggcagcc gcggtgacta taaatgaact caaagggaca ggtcatgtta  
 540

tgatgaaac caactgggtc gatgttgaat ttctcaacac tgcaaaacca ccacttggg  
 600

gttatcctgc ctcaaaaatg ctagctgaaa aggctgcatg gaaatttgct gaagaaaatg  
 660

acattgatct aatcactgtg atacctagtt taacaactgg tccttctctc acaccagata  
 720

tcccatctag tgttggcttg gcaatgtctc taataacagg caatgatatt ctcataaatg  
 780

ctttgaaagg aatgcagttt ctgtcgggtt cgttatccat cactcatgtt gaggatattt  
 840  
 gccgagctca tatatttctt gcagagaaag aatcagcttc tggtagatac atttgctgtg  
 900  
 ctcacaatac tagtgttccc gagcttgcaa agtttctcaa caaacgatat cctcagtata  
 960  
 aagttccaac tgaatttgat gattgcccc acaaggcaaa gttgataatc tcttctgaaa  
 1020  
 agcttatcaa agaagggttc agtttcaagc atggtattgc cgaaactttc gaccagactg  
 1080  
 tcgagtattt taagactaag ggggcactga agaattagat tttgatattt ctaattcaat  
 1140  
 agcaaactct aagcttggtt tgtgtttgtg aagttcagag tgaaatatca aatgaataag  
 1200  
 tggagagagc acaataagag gagagcacia taattttgga aaaaaaaaaa aaaaaaaaaa  
 1260  
 aaaaaaaaaagt actctgcgtt gttaccactg cttaatcact agtgaattc  
 1309

<210> 306  
 <211> 338  
 <212> PRT  
 <213> *Trifolium repens*

<400> 306

Met Ala Ser Ile Lys Gln Ile Gly Asn Lys Lys Ala Cys Val Ile Gly  
 1 5 10 15

Gly Thr Gly Phe Val Ala Ser Met Leu Ile Lys Gln Leu Leu Glu Lys  
 20 25 30

Gly Tyr Ala Val Asn Thr Thr Val Arg Asp Pro Asp Ser Pro Lys Lys  
 35 40 45

Ile Ser His Leu Val Ala Leu Gln Ser Leu Gly Glu Leu Asn Leu Phe  
 50 55 60

Arg Ala Asp Leu Thr Val Glu Glu Asp Phe Asp Ala Pro Ile Ala Gly  
 65 70 75 80

Cys Glu Leu Val Phe Gln Leu Ala Thr Pro Val Asn Phe Ala Ser Gln  
 85 90 95

Asp Pro Glu Asn Asp Met Ile Lys Pro Ala Ile Lys Gly Val Leu Asn  
 100 105 110

Val Leu Lys Ala Ile Ala Arg Ala Lys Glu Val Lys Arg Val Ile Leu  
 115 120 125

Thr Ser Ser Ala Ala Ala Val Thr Ile Asn Glu Leu Lys Gly Thr Gly  
 130 135 140  
 His Val Met Asp Glu Thr Asn Trp Ser Asp Val Glu Phe Leu Asn Thr  
 145 150 155 160  
 Ala Lys Pro Pro Thr Trp Gly Tyr Pro Ala Ser Lys Met Leu Ala Glu  
 165 170 175  
 Lys Ala Ala Trp Lys Phe Ala Glu Glu Asn Asp Ile Asp Leu Ile Thr  
 180 185 190  
 Val Ile Pro Ser Leu Thr Thr Gly Pro Ser Leu Thr Pro Asp Ile Pro  
 195 200 205  
 Ser Ser Val Gly Leu Ala Met Ser Leu Ile Thr Gly Asn Asp Phe Leu  
 210 215 220  
 Ile Asn Ala Leu Lys Gly Met Gln Phe Leu Ser Gly Ser Leu Ser Ile  
 225 230 235 240  
 Thr His Val Glu Asp Ile Cys Arg Ala His Ile Phe Leu Ala Glu Lys  
 245 250 255  
 Glu Ser Ala Ser Gly Arg Tyr Ile Cys Cys Ala His Asn Thr Ser Val  
 260 265 270  
 Pro Glu Leu Ala Lys Phe Leu Asn Lys Arg Tyr Pro Gln Tyr Lys Val  
 275 280 285  
 Pro Thr Glu Phe Asp Asp Cys Pro Ser Lys Ala Lys Leu Ile Ile Ser  
 290 295 300  
 Ser Glu Lys Leu Ile Lys Glu Gly Phe Ser Phe Lys His Gly Ile Ala  
 305 310 315 320  
 Glu Thr Phe Asp Gln Thr Val Glu Tyr Phe Lys Thr Lys Gly Ala Leu  
 325 330 335

Lys Asn

<210> 307

<211> 1005

<212> DNA

<213> Trifolium repens

<400> 307

gaattcgatt aagcagtggt aacaacgcag agtacgcggg gacttaaaca ttgacacaag  
60

tcccaaataa aaaagatctg aaacaacata gtcaccccat tttttaacat taaactaaaa  
120

atatgtcggc catcacgcga atccaagtcg agaacttga atttcggct gtggttactt  
180

ctcggccac cggtaagtca tattttcttg gtggtgcagg ggagagaggt ttgactattg  
240

aaggaaactt catcaagttc actgccatag gagtatatat ggaagatgta gcagtggctt  
300

catttgccac taaatggaag ggtaaactct ctgaggagtt gcttgagact cttgacttct  
360

atagagacat catttcagga ccctttgaaa agttgattcg aggatcgaag attaggggaat  
420

tgagtgggtc tgagtactca aggaagggtta atgaaaactg cgtggcacac ttaaaatctg  
480

ttgggactta tggagatgct gaagctgaag ctatgcaaaa atttggtgaa gccttcaagc  
540

ctattaatth tccacctggt gcctctgttt ttacaggca atcacctgat ggaatattag  
600

ggcttagttt ctctcaagat gcaagtatac cagaaaagga ggctgcagta atagagaaca  
660

aggcagcttc atcggcagtg ttagaaacta tgattggtga acatgctgtt tctcctgatt  
720

taaagcgttg tttggctgca agattacctg ccttggtgaa cgagggtact ttcaagattg  
780

aatgaaaact gattattatt atctccaaaa gcattgcagc acaagattga gtcatttatg  
840

agcatggaca tttttatgtc cacacatggt taacttttgt atctctcttt agattctcat  
900

caatatcaat aatactaata tgaaacgaag tcaaaaaaaaa aaaaaaaaaa aaaaaaaaaa  
960

aaaagtactc tgcgttggtta ccactgctta atcactagtg aattc  
1005

<210> 308

<211> 220

<212> PRT

<213> Trifolium repens

<400> 308

Met Ser Ala Ile Thr Ala Ile Gln Val Glu Asn Leu Glu Phe Pro Ala  
1 5 10 15

Val Val Thr Ser Pro Ala Thr Gly Lys Ser Tyr Phe Leu Gly Gly Ala  
20 25 30

Gly Glu Arg Gly Leu Thr Ile Glu Gly Asn Phe Ile Lys Phe Thr Ala  
35 40 45

Ile Gly Val Tyr Leu Glu Asp Val Ala Val Ala Ser Leu Ala Thr Lys  
 50 55 60

Trp Lys Gly Lys Ser Ser Glu Glu Leu Leu Glu Thr Leu Asp Phe Tyr  
 65 70 75 80

Arg Asp Ile Ile Ser Gly Pro Phe Glu Lys Leu Ile Arg Gly Ser Lys  
 85 90 95

Ile Arg Glu Leu Ser Gly Pro Glu Tyr Ser Arg Lys Val Asn Glu Asn  
 100 105 110

Cys Val Ala His Leu Lys Ser Val Gly Thr Tyr Gly Asp Ala Glu Ala  
 115 120 125

Glu Ala Met Gln Lys Phe Val Glu Ala Phe Lys Pro Ile Asn Phe Pro  
 130 135 140

Pro Gly Ala Ser Val Phe Tyr Arg Gln Ser Pro Asp Gly Ile Leu Gly  
 145 150 155 160

Leu Ser Phe Ser Gln Asp Ala Ser Ile Pro Glu Lys Glu Ala Ala Val  
 165 170 175

Ile Glu Asn Lys Ala Ala Ser Ser Ala Val Leu Glu Thr Met Ile Gly  
 180 185 190

Glu His Ala Val Ser Pro Asp Leu Lys Arg Cys Leu Ala Ala Arg Leu  
 195 200 205

Pro Ala Leu Leu Asn Glu Gly Thr Phe Lys Ile Glu  
 210 215 220

<210> 309  
 <211> 1105  
 <212> DNA  
 <213> Trifolium repens

<400> 309  
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 60

caacaccttc tccattacca tctatcttct actaagttca acgagatcaa tggcacttcc  
 120

ttctgtcacc gctttgaata tcgagaacaa tctattocct cctaccgtca caccaccggg  
 180

atccaccaac aatttcttcc tcggcggtgc aggagagcgg ggtcttcaaa ttcaagacaa  
 240

atttgtcaaa ttcaccgcta ttggtgttta tctacaggac attgctgttc cttacctcgc  
 300

cactaaatgg aagggttaaga ctgctcaaga gctaacggaa actgttcctt tcttcaggga  
360

catcgttaca ggtccatttg agaaatttat gcaggtgaca atgatcttgc cattgactgg  
420

gcaacaatac tcagagaaag tgtcagaaaa ttgtgtagct atttggaagt ctcttgggat  
480

ttataccgac gaagaagcca aagcaattga gaagtttggt tctgtcttca aagatgaaac  
540

attcccacca ggctcctcta tcctttttcac agtattaccc aaaggattag gatcactaac  
600

gataagtttc tctaaagatg gatccattcc agagaccgag tctgcagtta tagagaataa  
660

gctactctca caagctgtgc ttgagtcgat gataggggog cacggtgtct cccctgcagc  
720

aaaacagagt ttggccacca ggttatccga gttattcaac gaggttggtg atgctagcaa  
780

ctgattatat caacaaaacg aaaatgaaag tcctttctgc aataaagacc aagcggaat  
840

tttatttttag gtgcactttg aaatgacctc tttggcgact ttttcttgta ctaataataa  
900

agagtgtggt tgtatcatgt tgtaatttta ttttagaaaa agtgaggtaa gaaaggagtc  
960

cttatgttta tttcaattat tgaaaaatta tttgcatgta taattgattt caactgatgt  
1020

tatttaatca cgttttttct aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa gtactctgcg  
1080

ttgttaccac tgcttaatcg aattc  
1105

<210> 310

<211> 224

<212> PRT

<213> Trifolium repens

<400> 310

Met	Ala	Leu	Pro	Ser	Val	Thr	Ala	Leu	Asn	Ile	Glu	Asn	Asn	Leu	Phe
1				5					10					15	

Pro	Pro	Thr	Val	Thr	Pro	Pro	Gly	Ser	Thr	Asn	Asn	Phe	Phe	Leu	Gly
			20					25					30		

Gly	Ala	Gly	Glu	Arg	Gly	Leu	Gln	Ile	Gln	Asp	Lys	Phe	Val	Lys	Phe
		35					40					45			

Thr	Ala	Ile	Gly	Val	Tyr	Leu	Gln	Asp	Ile	Ala	Val	Pro	Tyr	Leu	Ala
		50					55					60			

Thr Lys Trp Lys Gly Lys Thr Ala Gln Glu Leu Thr Glu Thr Val Pro  
65 70 75 80

Phe Phe Arg Asp Ile Val Thr Gly Pro Phe Glu Lys Phe Met Gln Val  
85 90 95

Thr Met Ile Leu Pro Leu Thr Gly Gln Gln Tyr Ser Glu Lys Val Ser  
100 105 110

Glu Asn Cys Val Ala Ile Trp Lys Ser Leu Gly Ile Tyr Thr Asp Glu  
115 120 125

Glu Ala Lys Ala Ile Glu Lys Phe Val Ser Val Phe Lys Asp Glu Thr  
130 135 140

Phe Pro Pro Gly Ser Ser Ile Leu Phe Thr Val Leu Pro Lys Gly Leu  
145 150 155 160

Gly Ser Leu Thr Ile Ser Phe Ser Lys Asp Gly Ser Ile Pro Glu Thr  
165 170 175

Glu Ser Ala Val Ile Glu Asn Lys Leu Leu Ser Gln Ala Val Leu Glu  
180 185 190

Ser Met Ile Gly Ala His Gly Val Ser Pro Ala Ala Lys Gln Ser Leu  
195 200 205

Ala Thr Arg Leu Ser Glu Leu Phe Asn Glu Val Gly Asp Ala Ser Asn  
210 215 220

<210> 311

<211> 1272

<212> DNA

<213> Trifolium repens

<400> 311

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60

tgtgtaacaa atttcttaac ttaaaacatt ttcaacccaa caaaaaaaaa caaagacaaa  
120

aacatgggta gtgttgaaat tccaacaaag gttcttacta acagttctag tcaagtgaaa  
180

atgcctgtgg ttggaatggg atcagcacct gatttcacat gtaagaaaga cacaaaagat  
240

gcaatcattg aagccatcaa acagggttat agacactttg atactgctgc tgcttatggc  
300

tcagaacaag ctcttggtga aggtttgaaa gaagcaattg aacttggtct tgtcactaga  
360

gaagaccttt ttgttacttc taaactttgg gtcactgaaa atcatcctca tcttggtggt



420

cctgctcttc aaaaatctct caagactctt caattggagt acttggactt gtatttgatc  
480

cattggccac ttagttctca gcctggaaag ttttcatttc caattgatgt ggcagatctc  
540

ttgccatttg atgtgaaggg tgtttgggaa tccatggaag aaggcttgaa acttggactc  
600

actaaagcta ttggtgtag taacttctct gtcaagaaac ttcaaaatct tgtctcagtt  
660

gccactgttc ttctgctgt caatcaagtg gagatgaacc ttgcatggca acaaaagaag  
720

cttagagaat tttgcaatgc aaatggaata gtgttaactg cattttcacc attgagaaaa  
780

ggtgcaagca ggggaccaa tgaagttatg gaaaatgata tgcttaaaga gattgcagat  
840

gctcatggaa agtctgttgc acaaatttca ttgagatggg tatatgaaca aggagtcact  
900

tttgttccca agagctatga taaggaaaga atgggtcaaa atttggctat ctttgattgg  
960

acattggcaa aagaagatca tgagaaaatt gatcaaatta agcagaaccg tttgatccct  
1020

ggaccaacca agccaggact cagtgcacta tgggatgatg aaatataaag tggaagatgt  
1080

taaaagtccc ttaagctcac tcaatatcta tctattgtgt actttttgca tttgggggtt  
1140

gaaattgagt cacccttggt tctgtatcga tttaaaattt aaataatcaa tttttcatta  
1200

caaaaaaaaa aaaaaaaaaa aaaaaaaaaa agtactctgc gttgttacca ctgcttaatc  
1260

actagtgaat tc  
1272

&lt;210&gt; 312

&lt;211&gt; 314

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 312

Met Gly Ser Val Glu Ile Pro Thr Lys Val Leu Thr Asn Ser Ser Ser  
1 5 10 15

Gln Val Lys Met Pro Val Val Gly Met Gly Ser Ala Pro Asp Phe Thr  
20 25 30

Cys Lys Lys Asp Thr Lys Asp Ala Ile Ile Glu Ala Ile Lys Gln Gly  
35 40 45

Tyr Arg His Phe Asp Thr Ala Ala Ala Tyr Gly Ser Glu Gln Ala Leu  
 50 55 60

Gly Glu Gly Leu Lys Glu Ala Ile Glu Leu Gly Leu Val Thr Arg Glu  
 65 70 75 80

Asp Leu Phe Val Thr Ser Lys Leu Trp Val Thr Glu Asn His Pro His  
 85 90 95

Leu Val Val Pro Ala Leu Gln Lys Ser Leu Lys Thr Leu Gln Leu Glu  
 100 105 110

Tyr Leu Asp Leu Tyr Leu Ile His Trp Pro Leu Ser Ser Gln Pro Gly  
 115 120 125

Lys Phe Ser Phe Pro Ile Asp Val Ala Asp Leu Leu Pro Phe Asp Val  
 130 135 140

Lys Gly Val Trp Glu Ser Met Glu Glu Gly Leu Lys Leu Gly Leu Thr  
 145 150 155 160

Lys Ala Ile Gly Val Ser Asn Phe Ser Val Lys Lys Leu Gln Asn Leu  
 165 170 175

Val Ser Val Ala Thr Val Leu Pro Ala Val Asn Gln Val Glu Met Asn  
 180 185 190

Leu Ala Trp Gln Gln Lys Lys Leu Arg Glu Phe Cys Asn Ala Asn Gly  
 195 200 205

Ile Val Leu Thr Ala Phe Ser Pro Leu Arg Lys Gly Ala Ser Arg Gly  
 210 215 220

Pro Asn Glu Val Met Glu Asn Asp Met Leu Lys Glu Ile Ala Asp Ala  
 225 230 235 240

His Gly Lys Ser Val Ala Gln Ile Ser Leu Arg Trp Leu Tyr Glu Gln  
 245 250 255

Gly Val Thr Phe Val Pro Lys Ser Tyr Asp Lys Glu Arg Met Gly Gln  
 260 265 270

Asn Leu Ala Ile Phe Asp Trp Thr Leu Ala Lys Glu Asp His Glu Lys  
 275 280 285

Ile Asp Gln Ile Lys Gln Asn Arg Leu Ile Pro Gly Pro Thr Lys Pro  
 290 295 300

Gly Leu Ser Asp Leu Trp Asp Asp Glu Ile

305

310

<210> 313  
<211> 1548  
<212> DNA  
<213> *Trifolium repens*

<400> 313  
gaattcgatt aagcagtggg atcaacgcag agtacgcggg gacaacaact ataacttcct  
60

gttattaacc aattgagttc aaattacata catagcagga actatactaa agatatcaac  
120

atgggttagtg tttctgaaat tcgcaaggct caaagggctg aaggccctgc aactattttg  
180

gccattggta ctgcaaatcc agcaaactgt gttgaccaga gtacatatcc tgattttctac  
240

ttcaaaatca ctaacagtga gcataagggt gagcttaaag agaaatttca gcgcattgtg  
300

gataaatcta tgatcaagag cagatacatg tatctaacag aagagatttt gaaagaaaat  
360

cctagtcttt gtgaatacat ggcaccttca ttggatgcta ggcaagacat ggtggtggtt  
420

gaggtacctt gacttgggaa ggaggctgca gtgaaagcta tcaaagaatg ggggtcaacca  
480

aagtcaaaga ttactcactt aatcttttgc accacaagtg gtggtgacat gcctggtgac  
540

gattaccaac tcacaaaact cttagggtctt cgcccatatg tgaagaggta catgatgtac  
600

caacaagggt gctttgcagg tgggacgggt cttcgtttgg ccaaggattt ggccgagaac  
660

aacaaagggt ctcgtgtgtt ggttggttgc tctgaagtaa ccgcagtcac attccgcggc  
720

cccagtgaca ctcatttgga cagtcttggt ggacaagcac tattcgagga tggagctgct  
780

gcactcattg ttggctcaga cccagtacca gaaattgaga agccaatatt tgagatgggt  
840

tggaccgcac agacaattgc tccagatagt gaagggtgca ttgatgggtc tcttcgtgaa  
900

gctggactaa catttcatct tcttaaagat gttcctggga ttgtctcaaa gaacattgat  
960

aaggcattgg ttgaggcatt ccaaccatta aacatctctg attacaattc aatcttttgg  
1020

attgctcatc cagggtgggtc tgcaattcta gaccaagttg agataaagtt gggcttaaaa  
1080

cctgaaaaaa tgaaggccac cagagatgta cttagtgaat atggtaacat gtcaagtgca  
1140

tgtgtattgt tcattcttaga tgagatgaga aagaaatcgg ctgaaaatgg acttaaaacc  
1200

acaggagaag gacttgactg ggggtgtgttg tttggatttg ggcccggact taccattgaa  
1260

actgttggtc tacatagtgt ggctatatga gaatgagaga cttgatttgt ttttattgta  
1320

ttgtattgta ttactttaaa tcttggttga acctccattt taagaataaa tatggagtgc  
1380

aatatggacc atcctgttaa aataatatat cgtaaatagc tattatttta gtgtctgttt  
1440

ctttttacta aactatttta ttttagtatt tgtttttgac caaaaaaaaa aaaaaaaaaa  
1500

aaaaaaaaagta ctctgcgttg ttaccactgc ttaatcacta gtgaattc  
1548

<210> 314

<211> 389

<212> PRT

<213> *Trifolium repens*

<400> 314

Met Val Ser Val Ser Glu Ile Arg Lys Ala Gln Arg Ala Glu Gly Pro  
1 5 10 15

Ala Thr Ile Leu Ala Ile Gly Thr Ala Asn Pro Ala Asn Arg Val Asp  
20 25 30

Gln Ser Thr Tyr Pro Asp Phe Tyr Phe Lys Ile Thr Asn Ser Glu His  
35 40 45

Lys Val Glu Leu Lys Glu Lys Phe Gln Arg Met Cys Asp Lys Ser Met  
50 55 60

Ile Lys Ser Arg Tyr Met Tyr Leu Thr Glu Glu Ile Leu Lys Glu Asn  
65 70 75 80

Pro Ser Leu Cys Glu Tyr Met Ala Pro Ser Leu Asp Ala Arg Gln Asp  
85 90 95

Met Val Val Val Glu Val Pro Arg Leu Gly Lys Glu Ala Ala Val Lys  
100 105 110

Ala Ile Lys Glu Trp Gly Gln Pro Lys Ser Lys Ile Thr His Leu Ile  
115 120 125

Phe Cys Thr Thr Ser Gly Val Asp Met Pro Gly Ala Asp Tyr Gln Leu  
130 135 140

Thr Lys Leu Leu Gly Leu Arg Pro Tyr Val Lys Arg Tyr Met Met Tyr

145		150		155		160
Gln Gln Gly Cys Phe Ala Gly Gly Thr Val Leu Arg Leu Ala Lys Asp	165		170		175	
Leu Ala Glu Asn Asn Lys Gly Ala Arg Val Leu Val Val Cys Ser Glu	180		185		190	
Val Thr Ala Val Thr Phe Arg Gly Pro Ser Asp Thr His Leu Asp Ser	195		200		205	
Leu Val Gly Gln Ala Leu Phe Gly Asp Gly Ala Ala Ala Leu Ile Val	210		215		220	
Gly Ser Asp Pro Val Pro Glu Ile Glu Lys Pro Ile Phe Glu Met Val	225		230		235	240
Trp Thr Ala Gln Thr Ile Ala Pro Asp Ser Glu Gly Ala Ile Asp Gly		245		250		255
His Leu Arg Glu Ala Gly Leu Thr Phe His Leu Leu Lys Asp Val Pro		260		265		270
Gly Ile Val Ser Lys Asn Ile Asp Lys Ala Leu Val Glu Ala Phe Gln		275		280		285
Pro Leu Asn Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His Pro		290		295		300
Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Ile Lys Leu Gly Leu Lys		305		310		315
Pro Glu Lys Met Lys Ala Thr Arg Asp Val Leu Ser Glu Tyr Gly Asn		325		330		335
Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Lys Lys		340		345		350
Ser Ala Glu Asn Gly Leu Lys Thr Thr Gly Glu Gly Leu Asp Trp Gly		355		360		365
Val Leu Phe Gly Phe Gly Pro Gly Leu Thr Ile Glu Thr Val Val Leu		370		375		380
His Ser Val Ala Ile		385				

<210> 315  
 <211> 1447  
 <212> DNA

<213> Trifolium repens

<400> 315

gaattcacta gtgattaagc agtggttaaca acgcagagta cgcggggaac aaaaacaact  
60

acgcatatta tatatatata tatatagtct ataattgaaa gaaactgcta aagatattat  
120

taagatatgg tgagtgtagc tgaaattcgc aaggctcaga gggctgaagg ccctgcaacc  
180

attttgcca ttggcactgc aaatccacca aaccgtgttg agcagagcac atatcctgat  
240

ttctacttca aaattacaaa cagtgagcac aagactgagc tcaaagagaa gttccaacgc  
300

atgtgtgaca aatccatgat caagagcaga tacatgtatc taacagaaga gattttgaaa  
360

gaaaatccta gtctttgtga atacatggca ccttcattgg atgctaggca agacatgggt  
420

gtggttgagg tacctagact tgggaaggag gctgcagtca aggccattaa agaatggggg  
480

caaccaaagt caaagattac tcaacttaatc ttttgcacca caagtgggtg tgacatgcct  
540

ggtgctgatt accaactcac aaaactctta ggtcttgcgc catatgtgaa aaggtatatg  
600

atgtaccaac aagggtgttt tgcaggaggc acgggtgcttc gtttggcaaa agatttggcc  
660

gagaacaaca aagggtgctcg tgtgctagtt gtttgttctg aagtcaccgc agtcacattt  
720

cgcggcccca gtgatactca cttggacagt cttggtggac aagcattggt tggagatgga  
780

gccgctgcac taattgttgg ttctgatcca gtgcctgaaa ttgagaaacc aatatttgag  
840

atggtttgga ctgcacaaac aattgctcca gacagtgaag gtgccattga tggcatcctt  
900

cgtgaagctg ggctaacatt tcatottctt aaagatgttc ctgggattgt atcaaagaac  
960

attaataaag cattgggtga ggctttccaa ccattaggaa tttctgacta caactcaatc  
1020

ttttggattg cacaccggg tggacctgca attcttgatc aagtagaaca aaagctagcc  
1080

ttgaagcccg aaaagatgag ggccacgagg gaagttctaa gtgaatatgg aaacatgtca  
1140

agcgcagtgt tattgttcat cttagatgag atgcggaaga aatcggctca aaatggactt  
1200

aagacaactg gagaaggact tgattggggg gtgttggttc gcttcggacc aggacttacc  
1260

attgaaaccg ttgtttcttcg tagcgtggct atataagatg tgtgattggt tttatttttaa  
1320

tgtattactt ttaatcttgc tgccttgaat ttcgatttaa gaataaataa atatatcttt  
1380

tgataaaaaa aaaaaaaaaa aaaaaaaaaa aagtactctg cgttgttacc actgcttaat  
1440

cgaattc  
1447

<210> 316  
<211> 389  
<212> PRT  
<213> Trifolium repens  
  
<400> 316

Met Val Ser Val Ala Glu Ile Arg Lys Ala Gln Arg Ala Glu Gly Pro  
1 5 10 15

Ala Thr Ile Leu Ala Ile Gly Thr Ala Asn Pro Pro Asn Arg Val Glu  
20 25 30

Gln Ser Thr Tyr Pro Asp Phe Tyr Phe Lys Ile Thr Asn Ser Glu His  
35 40 45

Lys Thr Glu Leu Lys Glu Lys Phe Gln Arg Met Cys Asp Lys Ser Met  
50 55 60

Ile Lys Ser Arg Tyr Met Tyr Leu Thr Glu Glu Ile Leu Lys Glu Asn  
65 70 75 80

Pro Ser Leu Cys Glu Tyr Met Ala Pro Ser Leu Asp Ala Arg Gln Asp  
85 90 95

Met Val Val Val Glu Val Pro Arg Leu Gly Lys Glu Ala Ala Val Lys  
100 105 110

Ala Ile Lys Glu Trp Gly Gln Pro Lys Ser Lys Ile Thr His Leu Ile  
115 120 125

Phe Cys Thr Thr Ser Gly Val Asp Met Pro Gly Ala Asp Tyr Gln Leu  
130 135 140

Thr Lys Leu Leu Gly Leu Arg Pro Tyr Val Lys Arg Tyr Met Met Tyr  
145 150 155 160

Gln Gln Gly Cys Phe Ala Gly Gly Thr Val Leu Arg Leu Ala Lys Asp  
165 170 175

Leu Ala Glu Asn Asn Lys Gly Ala Arg Val Leu Val Val Cys Ser Glu  
180 185 190

Val Thr Ala Val Thr Phe Arg Gly Pro Ser Asp Thr His Leu Asp Ser  
195 200 205

Leu Val Gly Gln Ala Leu Phe Gly Asp Gly Ala Ala Ala Leu Ile Val  
210 215 220

Gly Ser Asp Pro Val Pro Glu Ile Glu Lys Pro Ile Phe Glu Met Val  
225 230 235 240

Trp Thr Ala Gln Thr Ile Ala Pro Asp Ser Glu Gly Ala Ile Asp Gly  
245 250 255

His Leu Arg Glu Ala Gly Leu Thr Phe His Leu Leu Lys Asp Val Pro  
260 265 270

Gly Ile Val Ser Lys Asn Ile Asn Lys Ala Leu Val Glu Ala Phe Gln  
275 280 285

Pro Leu Gly Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His Pro  
290 295 300

Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Gln Lys Leu Ala Leu Lys  
305 310 315 320

Pro Glu Lys Met Arg Ala Thr Arg Glu Val Leu Ser Glu Tyr Gly Asn  
325 330 335

Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Lys Lys  
340 345 350

Ser Ala Gln Asn Gly Leu Lys Thr Thr Gly Glu Gly Leu Asp Trp Gly  
355 360 365

Val Leu Phe Gly Phe Gly Pro Gly Leu Thr Ile Glu Thr Val Val Leu  
370 375 380

Arg Ser Val Ala Ile  
385

<210> 317  
<211> 2394  
<212> DNA  
<213> Trifolium repens

<400> 317  
gaattcgatt aagcagtggg aacaacgcag agtacgcggg gattcaatct gttgtgcata  
60

aaattcactc attgcataga aaaccataca catttgatct tgcaaagaag aaatatggga  
120



gacgaaggta tagtgagagg tgtcacaaag cagacaaccc ctgggaaggc tactatatattg  
180

gctcttggca aggcattccc tcaccaactt gtgatgcaag agtgtttagt tgatgggttat  
240

tttagggaca ctaattgtga caatcctgaa cttaagcaga aacttgctag actttgtaag  
300

acaaccacgg taaaaacaag gtatgttggt atgaatgagg agatactaaa gaaatatcca  
360

gaacttggtg tcgaaggcgc ctcaactgta aaacaacgtt tagagatatg taatgaggca  
420

gtaacacaaa tggcaattga agcttcccaa gtttgcctaa agaattgggg tagatcctta  
480

tcggacataa ctcatgtggt ttatgtttca tctagtgaag ctagattacc cggtggtgac  
540

ctatacttgt caaaaggact aggactaaac cctaaaattc aaagaaccat gctctatttc  
600

tctggatgct cgggaggcgt agccggcctt cgcgttgca aagacgtagc tgagaacaac  
660

cctggaagta gagttttgct tgctacttcg gaaactacaa ttattggatt caagccacca  
720

agtggtgata gaccttatga tcttggtggt gtggcactct ttggagatgg tgctggtgca  
780

atgataattg gctcagaccc ggtatttgaa actgagacac cattgtttga gctgcatact  
840

tcagctcagg agtttatacc agacaccgag aagaaaattg atgggcggct gacggaggag  
900

ggcataagtt tcacactagc aagggaactt ccgcagataa tcgaagacaa tgttgagggg  
960

ttctgtaata aactaattga tggtgttggg ttggagaata aggagtacaa taagttgttt  
1020

tgggctgtgc atccagggtg gcctgcgata ttgaatcgcg tggagaagcg gcttgagttg  
1080

tcgccgcaga agctgaatgc tagtagaaaa gctctaattg attatggaaa tgctagcagc  
1140

aatactattg tttatgtgct ggaatatatg ctagaagagg aaaagaagat taaaaaggcg  
1200

ggtgaggag attctgaatg gggattgata cttgcttttg gacctggaat tacttttgag  
1260

gggattctag caaggaactt gtgtgcatga agtcttatac aattgtgatg catgaactat  
1320

actcttattt ctactaatta ttatattaag caaattcaga acttttaagt aatgatttaa  
1380

tgaagaatac ttatagtata ttgactttat tcactttcaa agcaagttta tgatcctaag  
1440

acatggtaga acttgagcat gtggaatagt tgtaacaaaa actctaagca aatagagact  
 1500  
 ttatgtagta taaagcattt ccagacatga taaataatgg tacctcagaa cataaaatat  
 1560  
 atttagctat ctttcatccc caactttaca catccaccaa ggtacagaat aagcatatgt  
 1620  
 caacacaaaa tgtactctaa gtotaacatg agtaaccaa catgatgcct gattaagtta  
 1680  
 aaagaaaaga aaatctgagg gcatagatct tcaatcacac cactccagag ggaaggcgta  
 1740  
 gaacaagctg tccgccgaaa aactgcaat tcaataaata tcattaggac aacagtgcag  
 1800  
 agtcatgcgg gaaatgtctt aagtcaactgt actaaaaata taggattata ttatgaacta  
 1860  
 tactaacctt ttcacataat agtaacagaa atcagctaag atgaatgtct ggacaatttc  
 1920  
 tgagataaga accatgacgg ccataagcca taccccaagg caaccaataa atgtccacgg  
 1980  
 gtatctaaca cctggtgcaa gaaatagtaa gttattagga gatgtgcggg tacgaaattc  
 2040  
 aagctacaca acaaaaggag gccagaacaa cagcaatctt gtaaccagat gacaacaata  
 2100  
 aaatgtaaac ttaaagagac cgaacacaca aacattgcaa ctcagatgga attgctgcca  
 2160  
 tgtaactagt aggagatttg ggacgtcaaa tcagtatatt atgcaaatac aaggatatgac  
 2220  
 cgccttgtct attgtagcat acaacaaacg tacagtgggt ttgtccctct caaaatggca  
 2280  
 ggatctttac agcacaatat ttgggtttgt catacttata ccataaaaaa aaaaaaaaaa  
 2340  
 aaaaaaaaaa aaagtactct gcgttggtac cactgcttaa tcactagtga attc  
 2394

<210> 318  
 <211> 391  
 <212> PRT  
 <213> Trifolium repens  
 <400> 318

Met Gly Asp Glu Gly Ile Val Arg Gly Val Thr Lys Gln Thr Thr Pro  
 1 5 10 15

Gly Lys Ala Thr Ile Leu Ala Leu Gly Lys Ala Phe Pro His Gln Leu  
 20 25 30

Val Met Gln Glu Cys Leu Val Asp Gly Tyr Phe Arg Asp Thr Asn Cys  
 35 40 45

Asp Asn Pro Glu Leu Lys Gln Lys Leu Ala Arg Leu Cys Lys Thr Thr  
 50 55 60

Thr Val Lys Thr Arg Tyr Val Val Met Asn Glu Glu Ile Leu Lys Lys  
 65 70 75 80

Tyr Pro Glu Leu Val Val Glu Gly Ala Ser Thr Val Lys Gln Arg Leu  
 85 90 95

Glu Ile Cys Asn Glu Ala Val Thr Gln Met Ala Ile Glu Ala Ser Gln  
 100 105 110

Val Cys Leu Lys Asn Trp Gly Arg Ser Leu Ser Asp Ile Thr His Val  
 115 120 125

Val Tyr Val Ser Ser Ser Glu Ala Arg Leu Pro Gly Gly Asp Leu Tyr  
 130 135 140

Leu Ser Lys Gly Leu Gly Leu Asn Pro Lys Ile Gln Arg Thr Met Leu  
 145 150 155 160

Tyr Phe Ser Gly Cys Ser Gly Gly Val Ala Gly Leu Arg Val Ala Lys  
 165 170 175

Asp Val Ala Glu Asn Asn Pro Gly Ser Arg Val Leu Leu Ala Thr Ser  
 180 185 190

Glu Thr Thr Ile Ile Gly Phe Lys Pro Pro Ser Val Asp Arg Pro Tyr  
 195 200 205

Asp Leu Val Gly Val Ala Leu Phe Gly Asp Gly Ala Gly Ala Met Ile  
 210 215 220

Ile Gly Ser Asp Pro Val Phe Glu Thr Glu Thr Pro Leu Phe Glu Leu  
 225 230 235 240

His Thr Ser Ala Gln Glu Phe Ile Pro Asp Thr Glu Lys Lys Ile Asp  
 245 250 255

Gly Arg Leu Thr Glu Glu Gly Ile Ser Phe Thr Leu Ala Arg Glu Leu  
 260 265 270

Pro Gln Ile Ile Glu Asp Asn Val Glu Gly Phe Cys Asn Lys Leu Ile  
 275 280 285

Asp Val Val Gly Leu Glu Asn Lys Glu Tyr Asn Lys Leu Phe Trp Ala  
 290 295 300

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Val His Pro Gly Gly Pro Ala Ile Leu Asn Arg Val Glu Lys Arg Leu  
 305 310 315 320

Glu Leu Ser Pro Gln Lys Leu Asn Ala Ser Arg Lys Ala Leu Met Asp  
 325 330 335

Tyr Gly Asn Ala Ser Ser Asn Thr Ile Val Tyr Val Leu Glu Tyr Met  
 340 345 350

Leu Glu Glu Glu Lys Lys Ile Lys Lys Ala Gly Gly Gly Asp Ser Glu  
 355 360 365

Trp Gly Leu Ile Leu Ala Phe Gly Pro Gly Ile Thr Phe Glu Gly Ile  
 370 375 380

Leu Ala Arg Asn Leu Cys Ala  
 385 390

<210> 319  
 <211> 1663  
 <212> DNA  
 <213> Trifolium repens

<400> 319  
 gaattcgatt aagcagtggt aacaacgcag agtacgcggg gatagcaaca cacactttga  
 60  
 tttcttttttg agtccttgct acgtggcttt accaaaaaac gttgctaagt catcaacctt  
 120  
 tccaattcct taatataacc tatcagtact caccatcttt tcttctctcc tgctaacttt  
 180  
 agactcagag aagatgggtga atgttaatga gatccgccag gcacagagag ctgaaggccc  
 240  
 tgccaccgtg ttggcaatcg gcactgcaac tcttccaaac tgtgtcgatc agagtacata  
 300  
 cccagactac tacttccgca tcacaaacag tgagcacaag acagagctca aagaaaaatt  
 360  
 ccagcgcgatg tgtgacaaat ctatgattaa gaagagatac atgcatttga cagaagagat  
 420  
 tttgaaggag aatccaagtt tatgtgagta catggcacct tcattggatg caagacaaga  
 480  
 catgggtggtt gtggaagtac caaggctagg aaaagaggct gcaacaaagg ctatcaagga  
 540  
 atgggggtcaa cctaagtcca agattactca cctcatcttt tgcaaccacaa gtgggtgtgga  
 600  
 catgcctggc gccgactatc agcttacaaa gcttttaggc cttcgtccgc atgtgaagcg  
 660  
 ttatatgatg taccaacaag gttgtttcgc tgggtggtacg gtgcttcggt tggctaaaga  
 720

360/390

cttggctgaa aacaacaaag gtgcccggtg gttgggtggtt tgttcagaga tcaactgcggt  
780

tactttccgt ggacccagtg acactcatct tgatagcctt gtggggcaag cattgtttgg  
840

agatgggtgca gcagctgtga ttgtagggtc agaccatta ccacaagttg agaagccctt  
900

gtttgaattg gtatggactg ctcaaacaat ccttccagac agtgaaggag ccattgatgg  
960

gcaccttcgt gaagtcgggc tgacattcca tctcctcaag gatgttcctg gactcatctc  
1020

aaagaacatt gagaaagctc ttgttgaggc ctttcaacct ttaggtatct ctgattacaa  
1080

ttctatatatt tggatcgcac atcctggtgg acctgcaatt ctggaccaag tggaagccaa  
1140

attaagctta aagccagaga aaatgcaagc caccgcgcat gtgcttagcg agtatggtaa  
1200

catgtcaagt gcatgtgtgt tatttatctt ggatgagatg aggaggaagt caaaagaaga  
1260

tggacttgcc acaacaggcg aggggctgga atgggggtgta ctattcggtt ttggacccgg  
1320

actcactggt gagactgtat tgctccatag tgttgccact taaattgcct agatatgcta  
1380

taactatatg cttatttaat tctttgtttc tgggggattt tatcttcact tacttcactg  
1440

agcatttgaa taaagtttgt tttaattatt cataatgtaa tatgggtgttg cttaatgtac  
1500

ccatccatat aatatttgta atacatatat taatcaactt gcaatttcat gaaaaaaaaa  
1560

aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa aaggaaaaaa aaaaaaaaaa aaaaaaaaaa  
1620

aagtactctg cgttggtacc actgottaat cactagttaa ttc  
1663

<210> 320

<211> 389

<212> PRT

<213> Trifolium repens

<400> 320

Met Val Asn Val Asn Glu Ile Arg Gln Ala Gln Arg Ala Glu Gly Pro  
1 5 10 15

Ala Thr Val Leu Ala Ile Gly Thr Ala Thr Pro Pro Asn Cys Val Asp  
20 25 30

Gln Ser Thr Tyr Pro Asp Tyr Tyr Phe Arg Ile Thr Asn Ser Glu His  
35 40 45

Lys Thr Glu Leu Lys Glu Lys Phe Gln Arg Met Cys Asp Lys Ser Met  
 50 55 60  
 Ile Lys Lys Arg Tyr Met His Leu Thr Glu Glu Ile Leu Lys Glu Asn  
 65 70 75 80  
 Pro Ser Leu Cys Glu Tyr Met Ala Pro Ser Leu Asp Ala Arg Gln Asp  
 85 90 95  
 Met Val Val Val Glu Val Pro Arg Leu Gly Lys Glu Ala Ala Thr Lys  
 100 105 110  
 Ala Ile Lys Glu Trp Gly Gln Pro Lys Ser Lys Ile Thr His Leu Ile  
 115 120 125  
 Phe Cys Thr Thr Ser Gly Val Asp Met Pro Gly Ala Asp Tyr Gln Leu  
 130 135 140  
 Thr Lys Leu Leu Gly Leu Arg Pro His Val Lys Arg Tyr Met Met Tyr  
 145 150 155 160  
 Gln Gln Gly Cys Phe Ala Gly Gly Thr Val Leu Arg Leu Ala Lys Asp  
 165 170 175  
 Leu Ala Glu Asn Asn Lys Gly Ala Arg Val Leu Val Val Cys Ser Glu  
 180 185 190  
 Ile Thr Ala Val Thr Phe Arg Gly Pro Ser Asp Thr His Leu Asp Ser  
 195 200 205  
 Leu Val Gly Gln Ala Leu Phe Gly Asp Gly Ala Ala Ala Val Ile Val  
 210 215 220  
 Gly Ser Asp Pro Leu Pro Gln Val Glu Lys Pro Leu Phe Glu Leu Val  
 225 230 235 240  
 Trp Thr Ala Gln Thr Ile Leu Pro Asp Ser Glu Gly Ala Ile Asp Gly  
 245 250 255  
 His Leu Arg Glu Val Gly Leu Thr Phe His Leu Leu Lys Asp Val Pro  
 260 265 270  
 Gly Leu Ile Ser Lys Asn Ile Glu Lys Ala Leu Val Glu Ala Phe Gln  
 275 280 285  
 Pro Leu Gly Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His Pro  
 290 295 300

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Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Ala Lys Leu Ser Leu Lys  
 305 310 315 320

Pro Glu Lys Met Gln Ala Thr Arg His Val Leu Ser Glu Tyr Gly Asn  
 325 330 335

Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Arg Lys  
 340 345 350

Ser Lys Glu Asp Gly Leu Ala Thr Thr Gly Glu Gly Leu Glu Trp Gly  
 355 360 365

Val Leu Phe Gly Phe Gly Pro Gly Leu Thr Val Glu Thr Val Leu Leu  
 370 375 380

His Ser Val Ala Thr  
 385

<210> 321  
 <211> 1653  
 <212> DNA  
 <213> Trifolium repens

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 120

gatttgaatg gaagttcctc ggtgaatgga gcacgtgcta gacgtgctcc tactcaggga  
 180

aaggcaacga tacttgcatt aggaaaggct ttccccgccc aggtcctccc tcaagagtgc  
 240

ttggtggaag gattcattcg cgacactaag tgtgacgata cttatatataa ggagaaattg  
 300

gagcgtcttt gcaaaaacac aactgtgaaa acaagataca cagtaatgtc aaaggagatc  
 360

ttagacaact atccagagct agccatagat ggaacaccaa caataaggca aaagcttgaa  
 420

atagcaaadc cagcagtagt tgaaatggca acaagagcaa gcaaagattg catcaaagaa  
 480

tggggaaggc cacctcaaga tatcacacac atagtctatg tttcctcgag cgaaattcgt  
 540

ctaccgggtg gtgaccttta tcttgcaaag gaactcgggt taaacagcga tgttaatcgc  
 600

gtaatgctct atttcctcgg ttgctacggc ggtgtcactg gcttacgtgt cgccaaagac  
 660

atcgccgaaa ataaccctgg tagtaggggt ttactcacia catccgagac cactattctc  
 720

363/390

ggtttttcgac caccgagtaa agctagacct tatgacctcg ttggcgctgc acttttcggt  
780

gatggcgccg ctgctgcaat aattggaaca gacctatat tgaatcaaga atcacctttc  
840

atggaattga accatgcagt ccaaaaattc ttgcctgata cacaaaatgt gattgatggt  
900

agaatcactg aagagggtat taattttaag cttggaagag accttcctca aaaaattgaa  
960

gacaatattg aagaattttg caagaaaatt atggctaaaa gtgatgttaa ggaatttaat  
1020

gacttatttt gggctgttca tcttggtggg ccagctatac tcaataagct agaaaatata  
1080

ctcaaattga aaagtataa attggattgt agtaggaagg cattaatgga ttatggaaat  
1140

gttagtagca atactatatt ctatgtgatg gagtatatga gagattattt gaaggaagat  
1200

ggaagtgaag aatggggatt aggattggct tttggaccag ggattacttt tgaaggggtt  
1260

ctctccgta gcctttaatc ttgaaataat aattcatatg aaattacttg tcttaagatt  
1320

gtgataggaa gatgaatatg tattggatta atattgatat ggtgttattt taagttgatt  
1380

ttaaaaaaag tttattaata aagtatgatg taacaattgt tgtttgaatg ttaaaaggga  
1440

agtatactat ttttaagttct tgaccatact gattttttct ttacacattt tcatatctaa  
1500

aattgttcta tgatatcttc attgttgata ctgtaataat ataatatcta atttggtctg  
1560

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1620

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1653

<210> 322

<211> 389

<212> PRT

<213> Trifolium repens

<400> 322

Met Pro Gln Gly Asp Leu Asn Gly Ser Ser Ser Val Asn Gly Ala Arg  
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Ala Arg Arg Ala Pro Thr Gln Gly Lys Ala Thr Ile Leu Ala Leu Gly  
20 25 30

Lys Ala Phe Pro Ala Gln Val Leu Pro Gln Glu Cys Leu Val Glu Gly  
35 40 45



Phe Ile Arg Asp Thr Lys Cys Asp Asp Thr Tyr Ile Lys Glu Lys Leu  
 50 55 60

Glu Arg Leu Cys Lys Asn Thr Thr Val Lys Thr Arg Tyr Thr Val Met  
 65 70 75 80

Ser Lys Glu Ile Leu Asp Asn Tyr Pro Glu Leu Ala Ile Asp Gly Thr  
 85 90 95

Pro Thr Ile Arg Gln Lys Leu Glu Ile Ala Asn Pro Ala Val Val Glu  
 100 105 110

Met Ala Thr Arg Ala Ser Lys Asp Cys Ile Lys Glu Trp Gly Arg Ser  
 115 120 125

Pro Gln Asp Ile Thr His Ile Val Tyr Val Ser Ser Ser Glu Ile Arg  
 130 135 140

Leu Pro Gly Gly Asp Leu Tyr Leu Ala Asn Glu Leu Gly Leu Asn Ser  
 145 150 155 160

Asp Val Asn Arg Val Met Leu Tyr Phe Leu Gly Cys Tyr Gly Gly Val  
 165 170 175

Thr Gly Leu Arg Val Ala Lys Asp Ile Ala Glu Asn Asn Pro Gly Ser  
 180 185 190

Arg Val Leu Leu Thr Thr Ser Glu Thr Thr Ile Leu Gly Phe Arg Pro  
 195 200 205

Pro Ser Lys Ala Arg Pro Tyr Asp Leu Val Gly Ala Ala Leu Phe Gly  
 210 215 220

Asp Gly Ala Ala Ala Ala Ile Ile Gly Thr Asp Pro Ile Leu Asn Gln  
 225 230 235 240

Glu Ser Pro Phe Met Glu Leu Asn His Ala Val Gln Lys Phe Leu Pro  
 245 250 255

Asp Thr Gln Asn Val Ile Asp Gly Arg Ile Thr Glu Glu Gly Ile Asn  
 260 265 270

Phe Lys Leu Gly Arg Asp Leu Pro Gln Lys Ile Glu Asp Asn Ile Glu  
 275 280 285

Glu Phe Cys Lys Lys Ile Met Ala Lys Ser Asp Val Lys Glu Phe Asn  
 290 295 300

365/390

Asp Leu Phe Trp Ala Val His Pro Gly Gly Pro Ala Ile Leu Asn Lys  
 305 310 315 320

Leu Glu Asn Ile Leu Lys Leu Lys Ser Asp Lys Leu Asp Cys Ser Arg  
 325 330 335

Lys Ala Leu Met Asp Tyr Gly Asn Val Ser Ser Asn Thr Ile Phe Tyr  
 340 345 350

Val Met Glu Tyr Met Arg Asp Tyr Leu Lys Glu Asp Gly Ser Glu Glu  
 355 360 365

Trp Gly Leu Gly Leu Ala Phe Gly Pro Gly Ile Thr Phe Glu Gly Val  
 370 375 380

Leu Leu Arg Ser Leu  
 385

<210> 323  
 <211> 1600  
 <212> DNA  
 <213> Trifolium repens

<400> 323  
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 120

tatcttgggt acatcttttg ttacctccaa caaaaaaatg gtgaccgtag aagagattcg  
 180

taacgccccaa cgttcaaag ggcctgccac tatcttagct tttggcacag cactccttc  
 240

taactgtgtc actcaagctg attatcctga ttactacttt cgtatcacca acagcgaaca  
 300

tatgactgat ctttaaggaaa aattcaagcg gatgtgtgat agatcaatga taaagaaacg  
 360

ttacatgcac ctaacagaag actttctgaa ggagaatcca aatatgtgtg aatacatggc  
 420

accatcacta gatgtaagac gagacatagt ggttggtgaa gtaccaaagc taggtaaaga  
 480

agcagcaaaa aaagccatat gtgaatgggg acaacaaaaa tccaaaatca cacatcttgt  
 540

tttctgcacc acttcggtg ttgacatgcc gggagccgat taccaactca ccaaactttt  
 600

aggcttaaaa ccttctgtca agcgtctcat gatgtatcaa caagggtgtt tcgctggcgg  
 660

cacagttctc cgcttagcaa aagacottgt tgagaataac aaaaatgcaa gagttcttgt  
 720

366/390

tgtttgttct gaaattactg cggttacttt tcgtggacca tcggatactc atcttgattc  
780

gctcgtggga caggcgcttt ttggtgatgg agccgcagca atgattattg gtgcggatcc  
840

tgatttaacc gtggagcgtc cgattttcga gattgtttcg gctgctcaga ctattcttcc  
900

tgattctgat ggcgcaattg atggacatct tcgtgaagtg gggctcactt ttcatttatt  
960

gaaagatgtt ccgggggatta tttcaaagaa cattgaaaaa agtttagttg aagcttttgc  
1020

gcctattggg attaatgatt ggaactcaat attttgggtt gcacatccag gtggaccggc  
1080

tatttttagac caggttgaag agaaactcca tcttaaagag gagaaactcc ggtccaccgc  
1140

gcatgtgctt agtgaatatg gaaatatgtc aagtgcattg gttttattta ttttggatga  
1200

aatgagaaaag aggtctaaag aggaaggat gattacaact ggtgaagggt tggaatgggg  
1260

tgtgttgttt gggtttggac cgggtttaac tgttgaaacc gttgtgcttc atagtgttcc  
1320

ggttcagggt tgaatttatt atacatagat tggaaaataa aatttgcctg ccgagagatg  
1380

tgaactaact ttgtaggcaa gctcaaatta aagtttgaga taatattgtg ctttagttat  
1440

tatggtatgt aatgtaatgt ttttactttt ttcgaaattc atgtaatttg atatgtaaag  
1500

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1560

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<210> 324

<211> 391

<212> PRT

<213> Trifolium repens

<400> 324

Met Val Thr Val Glu Glu Ile Arg Asn Ala Gln Arg Ser Asn Gly Pro  
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Ala Thr Ile Leu Ala Phe Gly Thr Ala Thr Pro Ser Asn Cys Val Thr  
20 25 30

Gln Ala Asp Tyr Pro Asp Tyr Tyr Phe Arg Ile Thr Asn Ser Glu His  
35 40 45

Met Thr Asp Leu Lys Glu Lys Phe Lys Arg Met Cys Asp Arg Ser Met

60

Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Glu Lys Leu His Leu Lys  
305 310 315 320

Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Lys Arg  
340 345 350

Ser Lys Glu Glu Gly Met Ile Thr Thr Gly Glu Gly Leu Glu Trp Gly  
355 360 365

Val	Leu	Phe	Gly	Phe	Gly	Pro	Gly	Leu	Thr	Val	Glu	Thr	Val	Val	Leu
	370					375					380				

His Ser Val Pro Val Gln Gly  
385 390

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<210> 325
<211> 1333
<212> DNA
<213> Trifolium repens
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<400> 325
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120

tttgcgtcac cggcggcagc ggatgcatcg gttcatggct agtccatctc cttctcctcc  
180

gcggctacac tgttcacgcc accgtccaaa atctcaatga tgagaacgaa acgaagcatc  
240

tagaagctct cgaaggagca caaactaatc tccgtctctt ccagatcgat ctctttaact  
300

acgacacaat cctcgctgct gtccgcgggt gcgtcggaat ttccacctc gcttcacctt  
360

gcactgtaga caaagttcat gatcctcaga aggagctttt ggatcctgca attaaagga  
420

ctttgaatgt gcttactgca gctaaggaag taggggtgaa gcgtgtgggt gttacctcgt  
480

ctgtctcggc gattactcct agtcctgatt ggccttctga tgttgttaaa agagaggatt  
540

gttgactga tgttgaatat tgcaagaaaa aagagttgtg gtatcgttg tccaaaacat  
600

tggtgagaa agctgcgtgg gatttttcca aagaaaatgg tttggatgtt gttgtggtga  
660

atcccggtac tgtgatgggt cctgttattc caccacggca taatgcaagc atgctcatgc  
720

ttgtgagact tcttgaaggc tgcgctgaaa catttgaaga ctattttatg ggattgggtcc  
780

acttcaaaga tgtagcattg gcgcataatt tgggtgtatga gaacaaagaa gcatctggta  
840

gacatgtgtg tgttgaaact atctctcact acggtgattt tgtggcaaaa gttgctgaac  
900

tttatccaga atatagtgtt cctaggatgc agcgagatac gcaacctgga ttgttgagag  
960

cgaatgatgg atcaaagaag ctcatagatt tggggtttgga attcattcca atggagcaaa  
1020

ttatcaagga tgctgtagag agtttgaaga acaaaggatt catttcttga atgatgttac  
1080

tggtcttttg agaaccctat agttaccaga gtatagacta aataatatat aggtgatggg  
1140

tcagagaatg agtacttatg tcatgagttg tgtctgtata atatgttttc tcaattotta  
1200

tatgtttaat tgctaattgtt aacttcaata tttatcagcc agtattgttt ttttaataaa  
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1320

cttaatcgaa ttc  
1333

<210> 326  
<211> 320  
<212> PRT  
<213> Trifolium repens

<400> 326

Met Ser Lys Leu Val Cys Val Thr Gly Gly Ser Gly Cys Ile Gly Ser  
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Trp Leu Val His Leu Leu Leu Leu Arg Gly Tyr Thr Val His Ala Thr  
20 25 30

Val Gln Asn Leu Asn Asp Glu Asn Glu Thr Lys His Leu Glu Ala Leu  
35 40 45

Glu Gly Ala Gln Thr Asn Leu Arg Leu Phe Gln Ile Asp Leu Leu Asn  
50 55 60

Tyr Asp Thr Ile Leu Ala Ala Val Arg Gly Cys Val Gly Ile Phe His  
65 70 75 80

Leu Ala Ser Pro Cys Thr Val Asp Lys Val His Asp Pro Gln Lys Glu  
85 90 95

Leu Leu Asp Pro Ala Ile Lys Gly Thr Leu Asn Val Leu Thr Ala Ala  
100 105 110

370/390

Lys Glu Val Gly Val Lys Arg Val Val Val Thr Ser Ser Val Ser Ala  
 115 120 125

Ile Thr Pro Ser Pro Asp Trp Pro Ser Asp Val Val Lys Arg Glu Asp  
 130 135 140

Cys Trp Thr Asp Val Glu Tyr Cys Lys Lys Lys Glu Leu Trp Tyr Pro  
 145 150 155 160

Leu Ser Lys Thr Leu Ala Glu Lys Ala Ala Trp Asp Phe Ser Lys Glu  
 165 170 175

Asn Gly Leu Asp Val Val Val Val Asn Pro Gly Thr Val Met Gly Pro  
 180 185 190

Val Ile Pro Pro Arg His Asn Ala Ser Met Leu Met Leu Val Arg Leu  
 195 200 205

Leu Glu Gly Cys Ala Glu Thr Phe Glu Asp Tyr Phe Met Gly Leu Val  
 210 215 220

His Phe Lys Asp Val Ala Leu Ala His Ile Leu Val Tyr Glu Asn Lys  
 225 230 235 240

Glu Ala Ser Gly Arg His Val Cys Val Glu Thr Ile Ser His Tyr Gly  
 245 250 255

Asp Phe Val Ala Lys Val Ala Glu Leu Tyr Pro Glu Tyr Ser Val Pro  
 260 265 270

Arg Met Gln Arg Asp Thr Gln Pro Gly Leu Leu Arg Ala Asn Asp Gly  
 275 280 285

Ser Lys Lys Leu Ile Asp Leu Gly Leu Glu Phe Ile Pro Met Glu Gln  
 290 295 300

Ile Ile Lys Asp Ala Val Glu Ser Leu Lys Asn Lys Gly Phe Ile Ser  
 305 310 315 320

<210> 327  
 <211> 1470  
 <212> DNA  
 <213> Trifolium repens

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 120

ccatcatcaa acaatggcac cagccaaaac tctaagttat ctctcacaac aaaacactct  
 180

cgagtcaagt ttcgttaggg aagaagatga gcgtccaaaa gttgcctaca ataacttcag  
240

caacgagatt ccaatcattt ctcttgctgg aattgatgag gttgatggtc gtagaacaga  
300

gatatgtaac aagattgttg aagcttgtga gaattggggg atttttcagg ttgttgatca  
360

tggtgttgat acaaaaacttg tttctgagat gaccggtttt gctagagagt tttttgcttt  
420

gccaccggaa gagaagctcc ggtttgacat gtccgggtgg aaaaagggtg gtttcattgt  
480

ctctagtcac cttcaaggag aagcagtga ggattggaga gagctagtga catatttttc  
540

atacccaatt aaacaaagag attattcaag gtggccagac aagccagaag gatggaaaga  
600

ggtaacagaa aaatacagtg aaacctaata gaatttagct tgcaagctat tggaagtttt  
660

atcagaagca atgggttttag aaaaagaagc tctaacaaaa gcatgtgttg atatggatca  
720

aaaagttggt ataaattatt acccaaatg ccctgaacct gacctcacac ttggccttaa  
780

acgtcacact gaccctggca caattactct tttgcttcaa gatcaagttg gtggtcttca  
840

agctaccaaa gataatggta agacgtggat tacagttcaa ccagttgaag gtgcttttgt  
900

tggttaatctt ggagaccatg gtcactatct aagtaatgga cggttcaaaa atgctgacca  
960

tcaagcagtg gtgaattcga actacagccg tttatcaata gcaacatttc aaaatccagc  
1020

tccagatgca actgtgtacc ctttgaagat tagagatggg gaaaaatctg tgttggaaga  
1080

accaatcact tttgctgaaa tgtatagaag gaagatgacc aaagaccttg aaattgctag  
1140

gatgaagaag ttggctaagg aacaacaact tagggacttg gaggagaaca agactaaata  
1200

tgaggccaaa cctttgaatg agatctttgc ttaattaatt agtcttaatt taaataataa  
1260

attttagact taatttacat ataataattt taattttttg ttcaattaat ctatgtttaa  
1320

tttgtcgta ttgtccacgt gtattaagct gcttggttgt gtgtgccttg gagaataatc  
1380

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tgttaccact gcttaatcac tagtgaattc  
1470



<210> 328  
 <211> 366  
 <212> PRT  
 <213> Trifolium repens

<400> 328

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Glu Ser Ser Phe Val Arg Glu Glu Asp Glu Arg Pro Lys Val Ala Tyr  
 20 25 30

Asn Asn Phe Ser Asn Glu Ile Pro Ile Ile Ser Leu Ala Gly Ile Asp  
 35 40 45

Glu Val Asp Gly Arg Arg Thr Glu Ile Cys Asn Lys Ile Val Glu Ala  
 50 55 60

Cys Glu Asn Trp Gly Ile Phe Gln Val Val Asp His Gly Val Asp Thr  
 65 70 75 80

Lys Leu Val Ser Glu Met Thr Arg Phe Ala Arg Glu Phe Phe Ala Leu  
 85 90 95

Pro Pro Glu Glu Lys Leu Arg Phe Asp Met Ser Gly Gly Lys Lys Gly  
 100 105 110

Gly Phe Ile Val Ser Ser His Leu Gln Gly Glu Ala Val Lys Asp Trp  
 115 120 125

Arg Glu Leu Val Thr Tyr Phe Ser Tyr Pro Ile Lys Gln Arg Asp Tyr  
 130 135 140

Ser Arg Trp Pro Asp Lys Pro Glu Gly Trp Lys Glu Val Thr Glu Lys  
 145 150 155 160

Tyr Ser Glu Asn Leu Met Asn Leu Ala Cys Lys Leu Leu Glu Val Leu  
 165 170 175

Ser Glu Ala Met Gly Leu Glu Lys Glu Ala Leu Thr Lys Ala Cys Val  
 180 185 190

Asp Met Asp Gln Lys Val Val Ile Asn Tyr Tyr Pro Lys Cys Pro Glu  
 195 200 205

Pro Asp Leu Thr Leu Gly Leu Lys Arg His Thr Asp Pro Gly Thr Ile  
 210 215 220

Thr Leu Leu Leu Gln Asp Gln Val Gly Gly Leu Gln Ala Thr Lys Asp

225                                      230                                      235                                      240  
 Asn Gly Lys Thr Trp Ile Thr Val Gln Pro Val Glu Gly Ala Phe Val  
    245                                      250                                      255  
 Val Asn Leu Gly Asp His Gly His Tyr Leu Ser Asn Gly Arg Phe Lys  
    260                                      265                                      270  
 Asn Ala Asp His Gln Ala Val Val Asn Ser Asn Tyr Ser Arg Leu Ser  
    275                                      280                                      285  
 Ile Ala Thr Phe Gln Asn Pro Ala Pro Asp Ala Thr Val Tyr Pro Leu  
    290                                      295                                      300  
 Lys Ile Arg Asp Gly Glu Lys Ser Val Leu Glu Glu Pro Ile Thr Phe  
 305                                      310                                      315                                      320  
 Ala Glu Met Tyr Arg Arg Lys Met Thr Lys Asp Leu Glu Ile Ala Arg  
    325                                      330                                      335  
 Met Lys Lys Leu Ala Lys Glu Gln Gln Leu Arg Asp Leu Glu Glu Asn  
    340                                      345                                      350  
 Lys Thr Lys Tyr Glu Ala Lys Pro Leu Asn Glu Ile Phe Ala  
    355                                      360                                      365  
  
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 120  
 ctaccacatc acacaacata acaaattaag aaatatttat tactatatta agatatggaa  
 180  
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 240  
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 300  
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 360  
 ccggttgctc gtcttgggtg cgagacacta accattttct aggtggctgc cattgctgca  
 420  
 cagcatgggt caacgggtgga gctatcgga tctgctagag ccggcggtta ggcaagcagt  
 480  
 gactgggtta tggagagtat gaacaaagg accgacagct acggtgtccc aacaggggtc

540

ggcgctacct cgcaccgccg aaccaaacaa ggtgggtgctt tgcagaaaga gctcataagg  
600

tttttgaatg ctggaatatt tggaaatgga actgagtcaa gccacacact accacacaca  
660

gccacaagag ctgccatgct agtgagaatc aacacacttc tocaaggcta ttcaggaatt  
720

agatttgaaa tcttagaagc tatcaccaag cttcttaaca acaatgtcac cccatgttta  
780

ccgcttcgcy gtacaatcac agcttcagga gatttagtcc ctctttctta cattgctggt  
840

ttactaaccg gacgacaaaa ttccaaggct catggacott ctggagaagt acttaatgca  
900

aaacaagctt ttcaatcagc tggaaatgat gccgagttct ttgaattaca accaaaagaa  
960

ggccttgccc ttgttaacgg aaccgctggt gggtctggtt tagcttctat tggtcttttt  
1020

gaggctaata tattggcggg gttgtctgaa gttctatctg caattttcgc tgaagttatg  
1080

caaggggaagc ccgaatttac cgatcatttg acacataagt tgaaacatca cctgggtcaa  
1140

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1200

aagaagttgc atgagatgga tcctttacag aagccaaaac aagatagata tgcacttagg  
1260

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1320

gagagagaga tcaactctgt caatgacaat cctttgattg atgtttcaag gaacaaggct  
1380

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1440

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1500

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1560

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1620

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1680

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1740

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1800

tcgggtcaaaa ataccgtaag ccaagtggcc aaaaagacac taaccatagg tgtcaatgga

1860

gaacttcac cttcaagatt ttgtgaaaaa gacttattga aagtgggtga tagggaaacat  
1920

gtctttgcct acattgatga tccttgtagt gctacatacc cattgatgca aaaactcagg  
1980

caagtactag tggatcatgc attagttaat ggagaaagtg agaagaattht gaacacatca  
2040

atcttccaaa agattgcaac ttttgaggaa gagttgaaaa acctttgccca aaagagggtg  
2100

aaagtgaag gattgcatat gaaagtggaa attcaacaat tccaaacaag atcaatggat  
2160

gcagatctta tccactctac aattttgtga gaaaggagtt gggaactggt ttgctaactg  
2220

gagaaaatgt catttcaccg ggtgaagagt gtgacaaaact attcacagct atgtgtcaag  
2280

gaaaaatcat tgatcctctt cttgaatgct tgggagagtga gaacgggtgct cctcttccaa  
2340

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2400

tgcgagtgtg gtaataatga ttaggtgttt tgtgccttta atgaaaaaaaa aaaaaaaaaa  
2460

aaaaaaaaaa aaaagtactc tgcgttggtta ccaactgctta atcactagtg aattc  
2515

&lt;210&gt; 330

&lt;211&gt; 671

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 330

Met	Glu	Val	Val	Ala	Ala	Ala	Ile	Thr	Lys	Asn	Asn	Gly	Lys	Ile	Asp
1				5					10					15	

Ser	Phe	Cys	Leu	Asn	His	Ala	Asn	Ala	Asn	Asn	Met	Lys	Val	Asn	Gly
			20					25					30		

Ala	Asp	Pro	Leu	Asn	Trp	Gly	Val	Ala	Ala	Glu	Ala	Met	Lys	Gly	Ser
		35					40					45			

His	Leu	Asp	Glu	Val	Lys	Arg	Met	Val	Glu	Glu	Tyr	Arg	Lys	Pro	Val
	50					55					60				

Val	Arg	Leu	Gly	Gly	Glu	Thr	Leu	Thr	Ile	Ser	Gln	Val	Ala	Ala	Ile
65					70					75					80

Ala	Ala	His	Asp	Gly	Ala	Thr	Val	Glu	Leu	Ser	Glu	Ser	Ala	Arg	Ala
				85					90					95	

Gly Val Lys Ala Ser Ser Asp Trp Val Met Glu Ser Met Asn Lys Gly  
 100 105 110

Thr Asp Ser Tyr Gly Val Pro Thr Gly Phe Gly Ala Thr Ser His Arg  
 115 120 125

Arg Thr Lys Gln Gly Gly Ala Leu Gln Lys Glu Leu Ile Arg Phe Leu  
 130 135 140

Asn Ala Gly Ile Phe Gly Asn Gly Thr Glu Ser Ser His Thr Leu Pro  
 145 150 155 160

His Thr Ala Thr Arg Ala Ala Met Leu Val Arg Ile Asn Thr Leu Leu  
 165 170 175

Gln Gly Tyr Ser Gly Ile Arg Phe Glu Ile Leu Glu Ala Ile Thr Lys  
 180 185 190

Leu Leu Asn Asn Asn Val Thr Pro Cys Leu Pro Leu Arg Gly Thr Ile  
 195 200 205

Thr Ala Ser Gly Asp Leu Val Pro Leu Ser Tyr Ile Ala Gly Leu Leu  
 210 215 220

Thr Gly Arg Pro Asn Ser Lys Ala His Gly Pro Ser Gly Glu Val Leu  
 225 230 235 240

Asn Ala Lys Gln Ala Phe Gln Ser Ala Gly Ile Asp Ala Glu Phe Phe  
 245 250 255

Glu Leu Gln Pro Lys Glu Gly Leu Ala Leu Val Asn Gly Thr Ala Val  
 260 265 270

Gly Ser Gly Leu Ala Ser Ile Val Leu Phe Glu Ala Asn Ile Leu Ala  
 275 280 285

Val Leu Ser Glu Val Leu Ser Ala Ile Phe Ala Glu Val Met Gln Gly  
 290 295 300

Lys Pro Glu Phe Thr Asp His Leu Thr His Lys Leu Lys His His Pro  
 305 310 315 320

Gly Gln Ile Glu Ala Ala Ala Ile Met Glu His Ile Leu Asp Gly Ser  
 325 330 335

Ala Tyr Val Lys Asp Ala Lys Lys Leu His Glu Met Asp Pro Leu Gln  
 340 345 350

Lys Pro Lys Gln Asp Arg Tyr Ala Leu Arg Thr Ser Pro Gln Trp Leu

377/390

355		360		365
Gly Pro Leu Ile Glu Val Ile Arg Phe Ser Thr Lys Ser Ile Glu Arg				
370		375		380
Glu Ile Asn Ser Val Asn Asp Asn Pro Leu Ile Asp Val Ser Arg Asn				
385		390		395
Lys Ala Leu His Gly Gly Asn Phe Gln Gly Thr Pro Ile Gly Val Ser				
	405		410	415
Met Asp Asn Thr Arg Leu Ala Leu Ala Ser Ile Gly Lys Leu Met Phe				
	420		425	430
Ala Gln Phe Ser Glu Leu Val Asn Asp Phe Tyr Asn Asn Gly Leu Pro				
	435		440	445
Ser Asn Leu Ser Ala Ser Arg Asn Pro Ser Leu Asp Tyr Gly Phe Lys				
	450		455	460
Gly Ser Glu Ile Ala Met Ala Ser Tyr Cys Ser Glu Leu Gln Tyr Leu				
465		470		475
Ala Asn Pro Val Thr Thr His Val Gln Ser Ala Glu Gln His Asn Gln				
	485		490	495
Asp Val Asn Ser Leu Gly Leu Ile Ser Ser Arg Lys Thr Tyr Glu Ala				
	500		505	510
Ile Glu Ile Leu Gln Leu Met Ser Ser Thr Phe Leu Ile Ala Leu Cys				
	515		520	525
Gln Ala Ile Asp Leu Arg His Leu Glu Glu Asn Leu Lys Asn Ser Val				
	530		535	540
Lys Asn Thr Val Ser Gln Val Ala Lys Lys Thr Leu Thr Ile Gly Val				
545		550		555
Asn Gly Glu Leu His Pro Ser Arg Phe Cys Glu Lys Asp Leu Leu Lys				
	565		570	575
Val Val Asp Arg Glu His Val Phe Ala Tyr Ile Asp Asp Pro Cys Ser				
	580		585	590
Ala Thr Tyr Pro Leu Met Gln Lys Leu Arg Gln Val Leu Val Asp His				
	595		600	605
Ala Leu Val Asn Gly Glu Ser Glu Lys Asn Leu Asn Thr Ser Ile Phe				
	610		615	620

Gln Lys Ile Ala Thr Phe Glu Glu Glu Leu Lys Asn Leu Cys Gln Lys  
 625 630 635 640

Arg Leu Lys Val Gln Gly Leu His Met Lys Val Glu Ile Gln Gln Phe  
 645 650 655

Gln Thr Arg Ser Met Asp Ala Asp Leu Ile His Ser Thr Ile Leu  
 660 665 670

<210> 331  
 <211> 2667  
 <212> DNA  
 <213> Trifolium repens

<400> 331  
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ttccttattt cccacccaac acaacataac aaatacattt cctctcctct catcacaatt  
 120

attactttct acaccccccc ctctcaacta ttattaacta acataatgga gggaattacc  
 180

aatggccatg ctgaagcaac tttttgcgtg accaaaagtg ttggtgatcc actcaactgg  
 240

ggtgcagccg cggagtcgtt gatggggagt catttggtg aggtgaagcg tatggtggag  
 300

gaataccgta atccattggt taaaattggc ggcgagacgc ttaccattgc tcaggtggct  
 360

ggaattgctt ctcatgatag tgggtgtgagg gtggagctgt ctgagtcgc cagggccggc  
 420

gttaaggcga gtagtggttg ggtgatggac agcatgaaca atgggactga tagttatggt  
 480

gttaccactg gtttcggcgc cacctctcac cggagaacca agcaggggtg tgccttgcat  
 540

aaggagctaa ttaggttttt gaatgccgga atatttggca atggtacaga atctaactgt  
 600

acactaccac acacagcaac cagagctgca atgcttgtga gaatcaacac tcttcttcaa  
 660

ggatattctg gaattagatt tgaaattttg gaagctatca caaagcttct aaacaacaac  
 720

attaccccat gtttaccact tcgtggtaca atcacggctt ccggtgatct cgttcgctt  
 780

tcctacattg ccggtttggt aaccggtaga ccgaactcca aagccgttgg accctccgga  
 840

gaaattctca atgcaaaaaga agcttttcaa cttgccggca ttggttctga gttttttgaa  
 900

ttgcagccaa aagaaggtct tgctcttggt aatggtactg ctggttggtc tggtttagct  
 960

tctattgttc tgtttgaagc aaatgtacta gctgttttgt ctgaagttat gtcggcgatt  
1020

ttcgctgaag ttatgcaagg gaaaccagaa ttcactgac atttgactca taagttgaaa  
1080

catcacctg gtcaaattga agctgctgca attatggaac atattttgga tggaagtgt  
1140

tatgttaaag cagctaagaa attacacgaa accgatcctt tacaaaagcc gaaacaagat  
1200

cgttatgcac ttagaacttc acctcaatgg cttggtcctt tgattgaagt gataagattt  
1260

tcaactaagt caattgagag agaaattaac tctgtcaatg ataacccttt gattgatgtt  
1320

tcaaggaaca aggccattca cggtggtaat tttcaaggaa cacctattgg agtttcaatg  
1380

gataacacac gtttagctct tgcttcaatt ggtaaactca tgtttgctca attctctgaa  
1440

cttgtaatg atttttacaa caacgggtta ccttcgaatc ttactgctag taggaacca  
1500

agcttggtt acggtttcaa gggatcgga attgccatgg cttcttattg ttctgagtta  
1560

caatatcttg ctaatcctgt caccacccat gtccaaagt cggagcaaca caatcaagat  
1620

gttaactctt tgggtttgat ttcttcaaga aaaacaaatg aagctattga gatcctaaag  
1680

ctcatgtctt cgacatttct gattgcactt tgtcaagcaa ttgatttaag gcatttggag  
1740

gaaaatctga ggaacactgt caagaacacg gtaagccaag tagcgaagag aacactcacc  
1800

accggtgtta atggagaact tcatccttct agattttgtg agaaagattt gctcaaagtt  
1860

gttgataggg agtatgtatt tgcttatgtc gacgatcctt gtctagctac ataccctttg  
1920

atgcaaaagt tgagacaagt gcttgtggat catgcattgg taaatgctga tggagagaag  
1980

aatttgaaca catcaatctt tcaaaagatt gcaacttttg aggatgaatt gaaagctatc  
2040

ttgccaaagg aagttgaaag tacaagaact gcatatgaaa atggacaatg tggaatttca  
2100

aacaagatta aggaatgcag gtcttatcca ttgtacaagt ttgttagaga ggagtttagga  
2160

accgcgttgc taaccggaga aaaaacgata tcgctgggag aagagtgtga caaattgttc  
2220

acagctatgt gccaaagtaa aattgttgat cctcttttgg aatgccttgg agagtggaaat  
2280



ggtgctcctc taccaatatg ttaattagca gaattaatat gtttctttga gaagtgattt  
2340

ctttatatat ttgtagtata ctatagtagt tgcattgaga agcaattggg ttgtctataa  
2400

gcctatggaa aatggcaaaa caattttctg ctcaaagcat cgtttattaa gttttcctta  
2460

aagtgttaag gaacttttaa ttgtttttgt aatagaattt catttggttg ccacaacttt  
2520

gggtgcaa atcacgtgat acatgtggtg tttgatgtaa atggtgtttt ctcaattaat  
2580

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2640

taccactgct taatcactag tgaattc  
2667

<210> 332  
<211> 712  
<212> PRT  
<213> Trifolium repens

<400> 332

Met	Glu	Gly	Ile	Thr	Asn	Gly	His	Ala	Glu	Ala	Thr	Phe	Cys	Val	Thr
1				5					10					15	

Lys	Ser	Val	Gly	Asp	Pro	Leu	Asn	Trp	Gly	Ala	Ala	Ala	Glu	Ser	Leu
			20					25					30		

Met	Gly	Ser	His	Leu	Asp	Glu	Val	Lys	Arg	Met	Val	Glu	Glu	Tyr	Arg
		35					40					45			

Asn	Pro	Leu	Val	Lys	Ile	Gly	Gly	Glu	Thr	Leu	Thr	Ile	Ala	Gln	Val
	50					55					60				

Ala	Gly	Ile	Ala	Ser	His	Asp	Ser	Gly	Val	Arg	Val	Glu	Leu	Ser	Glu
65					70					75				80	

Ser	Ala	Arg	Ala	Gly	Val	Lys	Ala	Ser	Ser	Gly	Trp	Val	Met	Asp	Ser
				85					90					95	

Met	Asn	Asn	Gly	Thr	Asp	Ser	Tyr	Gly	Val	Thr	Thr	Gly	Phe	Gly	Ala
			100					105					110		

Thr	Ser	His	Arg	Arg	Thr	Lys	Gln	Gly	Gly	Ala	Leu	Gln	Lys	Glu	Leu
		115					120					125			

Ile	Arg	Phe	Leu	Asn	Ala	Gly	Ile	Phe	Gly	Asn	Gly	Thr	Glu	Ser	Asn
	130					135					140				

Cys Thr Leu Pro His Thr Ala Thr Arg Ala Ala Met Leu Val Arg Ile  
 145 150 155 160

Asn Thr Leu Leu Gln Gly Tyr Ser Gly Ile Arg Phe Glu Ile Leu Glu  
 165 170 175

Ala Ile Thr Lys Leu Leu Asn Asn Asn Ile Thr Pro Cys Leu Pro Leu  
 180 185 190

Arg Gly Thr Ile Thr Ala Ser Gly Asp Leu Val Pro Leu Ser Tyr Ile  
 195 200 205

Ala Gly Leu Leu Thr Gly Arg Pro Asn Ser Lys Ala Val Gly Pro Ser  
 210 215 220

Gly Glu Ile Leu Asn Ala Lys Glu Ala Phe Gln Leu Ala Gly Ile Gly  
 225 230 235 240

Ser Glu Phe Phe Glu Leu Gln Pro Lys Glu Gly Leu Ala Leu Val Asn  
 245 250 255

Gly Thr Ala Val Gly Ser Gly Leu Ala Ser Ile Val Leu Phe Glu Ala  
 260 265 270

Asn Val Leu Ala Val Leu Ser Glu Val Met Ser Ala Ile Phe Ala Glu  
 275 280 285

Val Met Gln Gly Lys Pro Glu Phe Thr Asp His Leu Thr His Lys Leu  
 290 295 300

Lys His His Pro Gly Gln Ile Glu Ala Ala Ala Ile Met Glu His Ile  
 305 310 315 320

Leu Asp Gly Ser Ala Tyr Val Lys Ala Ala Lys Lys Leu His Glu Thr  
 325 330 335

Asp Pro Leu Gln Lys Pro Lys Gln Asp Arg Tyr Ala Leu Arg Thr Ser  
 340 345 350

Pro Gln Trp Leu Gly Pro Leu Ile Glu Val Ile Arg Phe Ser Thr Lys  
 355 360 365

Ser Ile Glu Arg Glu Ile Asn Ser Val Asn Asp Asn Pro Leu Ile Asp  
 370 375 380

Val Ser Arg Asn Lys Ala Ile His Gly Gly Asn Phe Gln Gly Thr Pro  
 385 390 395 400

Ile Gly Val Ser Met Asp Asn Thr Arg Leu Ala Leu Ala Ser Ile Gly  
 405 410 415

Lys Leu Met Phe Ala Gln Phe Ser Glu Leu Val Asn Asp Phe Tyr Asn  
                   420                                  425                                  430

Asn Gly Leu Pro Ser Asn Leu Thr Ala Ser Arg Asn Pro Ser Leu Asp  
                   435                                  440                                  445

Tyr Gly Phe Lys Gly Ser Glu Ile Ala Met Ala Ser Tyr Cys Ser Glu  
                   450                                  455                                  460

Leu Gln Tyr Leu Ala Asn Pro Val Thr Thr His Val Gln Ser Ala Glu  
                   465                                  470                                  475                                  480

Gln His Asn Gln Asp Val Asn Ser Leu Gly Leu Ile Ser Ser Arg Lys  
                                   485                                  490                                  495

Thr Asn Glu Ala Ile Glu Ile Leu Lys Leu Met Ser Ser Thr Phe Leu  
                                   500                                  505                                  510

Ile Ala Leu Cys Gln Ala Ile Asp Leu Arg His Leu Glu Glu Asn Leu  
                   515                                  520                                  525

Arg Asn Thr Val Lys Asn Thr Val Ser Gln Val Ala Lys Arg Thr Leu  
                   530                                  535                                  540

Thr Thr Gly Val Asn Gly Glu Leu His Pro Ser Arg Phe Cys Glu Lys  
                   545                                  550                                  555                                  560

Asp Leu Leu Lys Val Val Asp Arg Glu Tyr Val Phe Ala Tyr Val Asp  
                                   565                                  570                                  575

Asp Pro Cys Leu Ala Thr Tyr Pro Leu Met Gln Lys Leu Arg Gln Val  
                                   580                                  585                                  590

Leu Val Asp His Ala Leu Val Asn Ala Asp Gly Glu Lys Asn Leu Asn  
                   595                                  600                                  605

Thr Ser Ile Phe Gln Lys Ile Ala Thr Phe Glu Asp Glu Leu Lys Ala  
                   610                                  615                                  620

Ile Leu Pro Lys Glu Val Glu Ser Thr Arg Thr Ala Tyr Glu Asn Gly  
                   625                                  630                                  635                                  640

Gln Cys Gly Ile Ser Asn Lys Ile Lys Glu Cys Arg Ser Tyr Pro Leu  
                                   645                                  650                                  655

Tyr Lys Phe Val Arg Glu Glu Leu Gly Thr Ala Leu Leu Thr Gly Glu  
                   660                                  665                                  670

Lys Thr Ile Ser Leu Gly Glu Glu Cys Asp Lys Leu Phe Thr Ala Met  
 675 680 685

Cys Gln Gly Lys Ile Val Asp Pro Leu Leu Glu Cys Leu Gly Glu Trp  
 690 695 700

Asn Gly Ala Pro Leu Pro Ile Cys  
 705 710

<210> 333  
 <211> 2664  
 <212> DNA  
 <213> *Trifolium repens*

<400> 333  
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 120  
 tcacaattat tacttttctac accctcctct caactattat taactagcat aatggaggga  
 180  
 attaccaatg gccatgctga aacaactttt tgcgtgacca aaagtgttgg tgatccactc  
 240  
 aactgggggtg cagccgcgga gtcgttgacg gggagtcatt tggatgaggt gaagcgtatg  
 300  
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 360  
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 420  
 gccggcggtta aggcgagtag tgattgggtg atggatagca tgaacaatgg gactgatagt  
 480  
 tacgggtgta ccaccggttt tggtgccacc tctcaccgga gaaccaagca gggtggtgct  
 540  
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 600  
 aactgtacac taccacacac agcaactaga gctgcaatgc ttgtgagaat caacactctt  
 660  
 cttcaagggt actctggtat tagatttgaa attttgaag ctatcacaaa gttccaaac  
 720  
 aacaacatta ccccatgttt accacttcgt ggtacaatca cggcttccgg tgatcttggt  
 780  
 ccgctttcct acattgccgg tttgttaacc ggaagacca actccaaagc agttggacct  
 840  
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 900  
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1020

gcgatttttcg ctgaagttat gcaaggga ccggaattta ccgatcattt gactcataag  
1080

ttgaaacatc accctgggtca aattgaagct gctgcaatta tggaacatat tttggatgga  
1140

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1200

caagatcggt atgcacttag aacttcacct caatggcttg gtcctttgat tgaagtgata  
1260

agattttcaa ccaaatcgat tgaaagagaa attaactcgg tcaacgacaa ccctttgatc  
1320

gatgtttcaa ggaacaaggc cattcatggt ggtaactttc aaggaacacc tattggagtt  
1380

tcaatggata acacacgttt agctcttgct tcaattggta aactcatggt tgctcaattc  
1440

tctgaacttg ttaatgattt ttacaacaac gggttgcctt cgtatcttac tgctagtagg  
1500

aaccgagct tggactatgg tttcaaggga tcggaaattg ccatggcttc gtattgttcc  
1560

gagttacaat atcttgctaa tcctgtcacc acccatgtcc aaagtgccga gcaacacaac  
1620

caagatgtta actctttggg tttgatttct tctagaaaaa caaatgaagc tattgagatt  
1680

ctcaagctca tgtcttccac tttcttgatt gcattatgtc aagcaatcga cttaaggcac  
1740

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1800

ctcaccaccg ggtcaacgg agaacttcat tcttctagat tttgtgagaa agatttgctt  
1860

aaagttgttg ataggagta tgtatttgcc tatgccgacg atccttgtct agctacatac  
1920

cctttgatgc aaaagttgag acaagtgcct gtggatcatg cattggtaaa tgttgatgga  
1980

gagaagaatt tgaacacatc aatctttcaa aagattgcaa cttttgagga tgagttgaaa  
2040

gctattttgc caaaggaggt tgaaagtaca agaactgcat atgaaaatgg acaatgtgga  
2100

atttcaaaca agattaagga atgcaggtct tatccattgt acaagtttgt tagagaggag  
2160

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2220

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2280

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2340

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2400

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2460

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2520

tttgggtgca aatatcacat gatacatgtg gtgtttgatg taaatggtgt tttttcaata  
2580

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2640

tgttaccact gcttaatoga attc  
2664

<210> 334  
<211> 712  
<212> PRT  
<213> Trifolium repens

<400> 334

Met Glu Gly Ile Thr Asn Gly His Ala Glu Thr Thr Phe Cys Val Thr  
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Lys Ser Val Gly Asp Pro Leu Asn Trp Gly Ala Ala Ala Glu Ser Leu  
20 25 30

Thr Gly Ser His Leu Asp Glu Val Lys Arg Met Val Glu Glu Tyr Arg  
35 40 45

Asn Pro Leu Ala Lys Ile Gly Gly Glu Thr Leu Thr Ile Ala Gln Val  
50 55 60

Ala Gly Ile Ala Ser His Asp Ser Gly Val Arg Val Glu Leu Ser Glu  
65 70 75 80

Ser Ala Arg Ala Gly Val Lys Ala Ser Ser Asp Trp Val Met Asp Ser  
85 90 95

Met Asn Asn Gly Thr Asp Ser Tyr Gly Val Thr Thr Gly Phe Gly Ala  
100 105 110

Thr Ser His Arg Arg Thr Lys Gln Gly Gly Ala Leu Gln Lys Glu Leu  
115 120 125

Ile Arg Phe Leu Asn Ala Gly Ile Phe Gly Asn Gly Thr Glu Ser Asn  
130 135 140

Cys Thr Leu Pro His Thr Ala Thr Arg Ala Ala Met Leu Val Arg Ile

145		150		155		160
Asn Thr Leu Leu Gln Gly Tyr Ser Gly Ile Arg Phe Glu Ile Leu Glu						
		165		170		175
Ala Ile Thr Lys Leu Pro Asn Asn Asn Ile Thr Pro Cys Leu Pro Leu						
		180		185		190
Arg Gly Thr Ile Thr Ala Ser Gly Asp Leu Val Pro Leu Ser Tyr Ile						
		195		200		205
Ala Gly Leu Leu Thr Gly Arg Pro Asn Ser Lys Ala Val Gly Pro Ser						
		210		215		220
Gly Glu Ile Leu Ser Ala Lys Glu Ala Phe Gln Leu Ala Gly Ile Gly						
		225		230		235
Ser Glu Phe Phe Glu Leu Gln Pro Lys Glu Gly Leu Ala Leu Val Asn						
		245		250		255
Gly Thr Ala Val Gly Ser Gly Leu Ala Ser Ile Val Leu Phe Glu Ala						
		260		265		270
Asn Val Leu Ala Val Leu Ser Glu Val Met Ser Ala Ile Phe Ala Glu						
		275		280		285
Val Met Gln Gly Lys Pro Glu Phe Thr Asp His Leu Thr His Lys Leu						
		290		295		300
Lys His His Pro Gly Gln Ile Glu Ala Ala Ala Ile Met Glu His Ile						
		305		310		315
Leu Asp Gly Ser Ala Tyr Val Lys Ala Ala Lys Lys Leu His Glu Thr						
		325		330		335
Asp Pro Leu Gln Lys Pro Lys Gln Asp Arg Tyr Ala Leu Arg Thr Ser						
		340		345		350
Pro Gln Trp Leu Gly Pro Leu Ile Glu Val Ile Arg Phe Ser Thr Lys						
		355		360		365
Ser Ile Glu Arg Glu Ile Asn Ser Val Asn Asp Asn Pro Leu Ile Asp						
		370		375		380
Val Ser Arg Asn Lys Ala Ile His Gly Gly Asn Phe Gln Gly Thr Pro						
		385		390		395
Ile Gly Val Ser Met Asp Asn Thr Arg Leu Ala Leu Ala Ser Ile Gly						
		405		410		415

Lys Leu Met Phe Ala Gln Phe Ser Glu Leu Val Asn Asp Phe Tyr Asn  
 420 425 430  
 Asn Gly Leu Pro Ser Tyr Leu Thr Ala Ser Arg Asn Pro Ser Leu Asp  
 435 440 445  
 Tyr Gly Phe Lys Gly Ser Glu Ile Ala Met Ala Ser Tyr Cys Ser Glu  
 450 455 460  
 Leu Gln Tyr Leu Ala Asn Pro Val Thr Thr His Val Gln Ser Ala Glu  
 465 470 475 480  
 Gln His Asn Gln Asp Val Asn Ser Leu Gly Leu Ile Ser Ser Arg Lys  
 485 490 495  
 Thr Asn Glu Ala Ile Glu Ile Leu Lys Leu Met Ser Ser Thr Phe Leu  
 500 505 510  
 Ile Ala Leu Cys Gln Ala Ile Asp Leu Arg His Leu Glu Glu Asn Leu  
 515 520 525  
 Arg Asn Thr Val Lys Asn Thr Val Ser Gln Val Ala Lys Arg Thr Leu  
 530 535 540  
 Thr Thr Gly Val Asn Gly Glu Leu His Ser Ser Arg Phe Cys Glu Lys  
 545 550 555 560  
 Asp Leu Leu Lys Val Val Asp Arg Glu Tyr Val Phe Ala Tyr Ala Asp  
 565 570 575  
 Asp Pro Cys Leu Ala Thr Tyr Pro Leu Met Gln Lys Leu Arg Gln Val  
 580 585 590  
 Leu Val Asp His Ala Leu Val Asn Val Asp Gly Glu Lys Asn Leu Asn  
 595 600 605  
 Thr Ser Ile Phe Gln Lys Ile Ala Thr Phe Glu Asp Glu Leu Lys Ala  
 610 615 620  
 Ile Leu Pro Lys Glu Val Glu Ser Thr Arg Thr Ala Tyr Glu Asn Gly  
 625 630 635 640  
 Gln Cys Gly Ile Ser Asn Lys Ile Lys Glu Cys Arg Ser Tyr Pro Leu  
 645 650 655  
 Tyr Lys Phe Val Arg Glu Glu Leu Gly Thr Ala Leu Leu Thr Gly Glu  
 660 665 670  
 Lys Thr Ile Ser Pro Gly Glu Glu Cys Asp Lys Leu Phe Thr Ala Met



675

680

685

Cys Gln Gly Lys Ile Val Asp Pro Leu Met Glu Cys Leu Gly Glu Trp  
690 695 700

Asn Gly Ala Pro Leu Pro Ile Cys  
705 710

<210> 335  
<211> 1296  
<212> DNA  
<213> Trifolium repens

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1200

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&lt;210&gt; 336

&lt;211&gt; 326

&lt;212&gt; PRT

<213> *Trifolium repens*

&lt;400&gt; 336

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Asn Thr Thr Ile Arg Ala Asp Pro Glu Arg Lys Arg Asp Val Ser Phe  
35 40 45

Leu Thr Asn Leu Pro Gly Ala Ser Glu Arg Leu His Phe Phe Asn Ala  
50 55 60

Asp Leu Asp Asp Pro Glu Ser Phe Asn Glu Ala Ile Glu Gly Cys Val  
65 70 75 80

Gly Ile Phe His Thr Ala Ser Pro Ile Asp Phe Ala Val Ser Glu Pro  
85 90 95

Glu Glu Ile Val Thr Lys Arg Thr Val Asp Gly Ala Leu Gly Ile Leu  
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Lys Ala Cys Val Asn Ser Lys Thr Val Lys Arg Phe Ile Tyr Thr Ser  
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130 135 140

Glu Ser Asp Trp Ser Asp Val Asp Leu Leu Arg Ser Val Lys Pro Phe  
145 150 155 160

Gly Trp Ser Tyr Gly Val Ser Lys Thr Leu Ala Glu Lys Ala Val Leu

Lys Glu Lys Gly Tyr Leu  
325

# INTERNATIONAL SEARCH REPORT

International application No.  
**PCT/AU02/01345**

<b>A. CLASSIFICATION OF SUBJECT MATTER</b>					
Int. Cl. <sup>7</sup> : C12N 15/29, 15/52, 15/53, 15/55, 15/60, 15/61; A01H 5/00					
According to International Patent Classification (IPC) or to both national classification and IPC					
<b>B. FIELDS SEARCHED</b>					
Minimum documentation searched (classification system followed by classification symbols) SEE ELECTRONIC DATABASE BOX BELOW					
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched SEE ELECTRONIC DATABASE BOX BELOW					
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) PEPTIDE DATABASES (SWISSPROT, GENBANK, EMBL, PIR) DGENE: SEQ ID NOS 2,9,14,18,24,65,70,79,92,96,109,111,118,136,148,154,156,160,162,164,169,186,195,197,203,246,248,250,279,287,29 4,299,304,308,310,312,314,318,320,322,324,326,328,334					
<b>C. DOCUMENTS CONSIDERED TO BE RELEVANT</b>					
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.			
PX	WO 02 18604 A (THE SALK INSTITUTE FOR BIOLOGICAL STUDIES) 7 March 2002 Fig 1D shares ~92% identity with SEQ ID NOS 2, 9, 14, 308	1-3, 13-19, 25, 27			
X	US 6 054 636 A (FADER GM) 25 April 2000 Fig 2 shares ~81% identity with SEQ ID NOS 2, 9, 14, 308	1-3, 13-27			
<input checked="" type="checkbox"/> Further documents are listed in the continuation of Box C <input checked="" type="checkbox"/> See patent family annex					
<table style="width: 100%; border: none;"> <tr> <td style="width: 33%; vertical-align: top;">           * Special categories of cited documents:            "A" document defining the general state of the art which is not considered to be of particular relevance            "E" earlier application or patent but published on or after the international filing date            "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)            "O" document referring to an oral disclosure, use, exhibition or other means            "P" document published prior to the international filing date but later than the priority date claimed         </td> <td style="width: 33%; vertical-align: top;">           "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention            "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone            "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art            "&amp;" document member of the same patent family         </td> <td style="width: 33%;"></td> </tr> </table>			* Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "&" document member of the same patent family	
* Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "&" document member of the same patent family				
Date of the actual completion of the international search 28 November 2002		Date of mailing of the international search report <b>05 DEC 2002</b>			
Name and mailing address of the ISA/AU AUSTRALIAN PATENT OFFICE PO BOX 200, WODEN ACT 2606, AUSTRALIA E-mail address: pct@ipaustalia.gov.au Facsimile No. (02) 6285 3929		Authorized officer  <b>TERRY MOORE</b> Telephone No : (02) 6283 2632			

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU02/01345

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 99 36543 A (PIONEER HI-BRED INTERNATIONAL, INC.) 22 July 1999 SEQ ID NO 2 shares ~81% identity with SEQ ID NOS 2, 9, 14, 308	1-3, 13-17, 21, 22, 24-27
X	Genbank Acc No AAB41524 chalcone isomerase ( <i>Medicago sativa</i> ) 29 January 1997 (See also Medline Abstract 8193301) 92% identity with SEQ ID NOS 18, 310	1-3, 13-18, 24-27
X	Genbank Acc No CAA74847 anther-specific protein ( <i>Nicotiana sylvestris</i> ) (See also Medline Abstract 99084767) 83% identity with SEQ ID NOS 96, 322 and 67% identity with SEQ ID NOS 318, 70	1, 2, 4, 13-18, 24-26, 28
X	Genbank Acc No CAC14061 chalcone synthase ( <i>Ruta graveolens</i> ) 27 October 2000 ~87% identity with SEQ ID NOS 24, 65, 79, 92, 102, 107, 314, 316, 320, 324	1, 2, 4, 13-18, 24-26, 28
X	Genbank Acc No AAB41556 chalcone reductase ( <i>Medicago sativa</i> ) 30 January 1997 ~95% identity with SEQ ID NOS 109, 118, 312	1, 2, 5, 13-18, 20, 24-26, 29
X	Genbank Acc No CAA11226 chalcone reductase ( <i>Sesbania rostrata</i> ) 3 July 2001 90% identity with SEQ ID NO 111	1, 2, 5, 13-18, 24-26, 29
X	Genbank Acc No AAK52955 dihydro-flavonoid reductase-like protein ( <i>Zea mays</i> ) 14 May 2001 69% identity with SEQ ID NOS 287, 160 and 53% identity with SEQ ID NO 148	1, 2, 6, 13-18, 24-26, 30
PX	WO 02 063021 A (PIONEER HI-BRED INTERNATIONAL, INC.) 15 August 2002 SEQ ID NO 1 shares 74% identity with SEQ ID NOS 148, 160, 287	1, 2, 6, 13-18, 24-26, 30
X	Genbank Acc No AAD54273 dihydroflavonol-4-reductase DFR1 ( <i>Glycine max</i> ) 10 September 1999 81 % identity with SEQ ID NO 169	1, 2, 6, 13-18, 24-26, 30
X	WO 95 27790 A (CENTRE NATIONAL DE LA RECHERCHE SCIENTIFIQUE) 19 October 1995 SEQ ID NO 1 shares 43% identity with SEQ ID NOS 136, 156, 326, 39% identity with SEQ ID NO 154 and 51% identity with SEQ ID NO 294.	1, 2, 6, 13-18, 24-26, 30
PX	WO 02 10210 A (BAYER AKTIENGESELLSCHAFT) 7 February 2002 SEQ ID NO 2329 shares 51% identity with SEQ ID NO 164, SEQ ID NO 636 shares 62% identity with SEQ ID NO 186, SEQ ID NO 1573 shares 49% identity with SEQ ID NOS 246, 248, SEQ ID NO 2091 shares 57% identity with SEQ ID NOS 304, 299	1, 2, 7, 9, 13- 17, 24-26, 31, 32
X	WO 99 14351 (E.I. DU PONT DE NEMOURS AND COMPANY) 25 March 1999 SEQ ID NO 1 shares 56% identity with SEQ ID NO 162	1, 2, 6, 13-17, 21, 22, 24-26, 30

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU02/01345

C (Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
PX	WO 02 26994 A (AGRICULTURE VICTORIA SERVICES PTY LTD) 4 April 2002 Fig 39 shares 72% identity with SEQ ID NO 294	1, 2, 6, 13-18, 21, 22, 24-26, 30
X	WO 97 12982 A (CENTRE NATIONAL DE LA RECHERCHE SCIENTIFIQUE) 10 April 1997 SEQ ID NO 5 shares 55% identity with SEQ ID NO 294	1, 2, 6, 13-17, 24-26, 30
X	Genbank Acc No CAA80265 flavonoid 3',5'-hydroxylase ( <i>Petunia x hybrida</i> ) 7 December 1993 70% identity with SEQ ID NO 197	1, 2, 8, 13-18, 24-26, 32
X	EP 1 033 405 (CERES INCORPORATED) 6 September 2000 SEQ ID NO 49742 shares 57% identity with SEQ ID NO 195	1, 2, 8, 13-18, 24-26, 32
X	Genbank Acc No AAF23859 DFR-like protein ( <i>Arabidopsis thaliana</i> ) 11 January 2000 61% identity with SEQ ID NO 186	1, 2, 7, 13-18, 24-26, 31
X	Genbank Acc No BAB01697 oxidase-like protein ( <i>Arabidopsis thaliana</i> ) 27 December 2000 50% identity with SEQ ID NOS 246, 248	1, 2, 9, 13-18, 24-26, 33
X	TREMBL Acc No CAB63776 F3'H1 protein ( <i>Glycine max</i> ) 1 May 2000 85% identity with SEQ ID NO 328, 203	1, 2, 9, 13-18, 24-26, 33
X	Genbank Acc No CAB78172 flavanone 3-beta-hydroxylase ( <i>Arabidopsis thaliana</i> ) 16 March 2000 57% identity with SEQ ID NOS 304, 299	1, 2, 9, 13-17, 24-26, 33
X	Genbank Acc No AAG49298 Flavonoid 3'-hydroxylase ( <i>Callistephus chinensis</i> ) 16 January 2001 68% identity with SEQ ID NO 250	1, 2, 10, 13- 18, 20, 24-26, 34
X	Genbank Acc No AAA99500 Phenylalanine ammonia lyase ( <i>Stylosanthes humilis</i> ) 15 May 1996 88% identity with SEQ ID NOS 254, 259, 269, 271, 273, 275, 330, 332, 334	1, 2, 11, 13- 18, 24-26, 35
X	Genbank Acc No CAA41169 phenylalanine ammonia lyase ( <i>Medicago sativa</i> ) 5 May 1995 87% identity with SEQ ID NOS 254, 259, 269, 271, 273, 275, 330, 332, 334	1, 2, 11, 13- 18, 24-26, 35
X	Genbank Acc No AAB41550 vestitone reductase ( <i>Medicago sativa</i> ) 30 January 1997 (See also Medline abstract 7625843) 95% identity with SEQ ID NOS 336, 279	1, 2, 12-18, 21, 22, 24-26, 36

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU02/01345

### Box I Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claims Nos :  
because they relate to subject matter not required to be searched by this Authority, namely:
2. ☐ Claims Nos :  
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. ☐ Claims Nos :  
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)

### Box II Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

See supplemental Box

1. ☒ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims
2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

- ☐ The additional search fees were accompanied by the applicant's protest.
- ☒ No protest accompanied the payment of additional search fees.

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU02/01345

**Supplemental Box 1**

(To be used when the space in any of Boxes I to VIII is not sufficient)

**Continuation of Box No: II (lack of unity)**

The international application does not comply with the requirements of unity of invention because it does not relate to one invention or to a group of inventions so linked as to form a single general inventive concept. The fundamental test for unity of invention is specified in Rule 13.2 of the Regulations under the PCT.

"Where a group of inventions is claimed in one and the same international application, the requirement of unity of invention referred to in Rule 13.1 shall be fulfilled only where there is a technical relationship among those inventions involving one or more of the same or corresponding special technical features. The expression "special technical feature" shall mean those technical features that define a contribution which each of the claimed inventions, considered as a whole, make over the prior art."

The problem addressed by the application is the modification of flavonoid biosynthesis (see page 2, line 29-page 3, line 7). The solution provided by the claims resides in the use of 56 specific polypeptides from clover, medic, ryegrass or fescue species (claim 25) and the nucleic acids or fragments coding for these polypeptides (claim 1). These 56 specific polypeptides fall within the following ten groups:

- |     |                                    |   |
|-----|------------------------------------|---|
| 1.  | Chalcone isomerase (CHI)           | (SEQ IDS 2,9,14,18,308,310)                             |
| 2.  | Chalcone synthase (CHS)            | (SEQ IDS 24,65,70,79,92,96,102,107,314,316,318,322,324) |
| 3.  | Chalcone reductase (CHR)           | (SEQ IDS 109,111,118,312)                               |
| 4.  | Dihydroflavonol 4-reductase (DFR)  | (SEQ IDS 136,148,154,156,160,162,164,169,287,294,326)   |
| 5.  | Leucoanthocyanidin reductase (LCR) | (SEQ ID 186)  |
| 6.  | Flavonoid 3',5' hydrolase (F3'5'H) | (SEQ IDS 195,197)                                       |
| 7.  | Flavanone 3-hydrolase (F3H)        | (SEQ IDS 203,246,248,299,304,328)                       |
| 8.  | Flavonoid 3'-hydroxylase (F3'H)    | (SEQ ID 250)  |
| 9.  | Phenylalanine ammonia-lyase (PAL)  | (SEQ IDS 254,259,269,271,273,275,330,332,334)           |
| 10. | Vestitone reductase (VR)           | (SEQ IDS 279,336)                                       |

The application acknowledges that representatives of these enzyme species, and the nucleotides that encode them, are known and have been isolated from other plant species (see page 2 lines 29-31). This is supported by the following documents, which disclose the isolation and characterisation of a number of these enzymes from a range of species, including clover and medic and rye.

EMBL CAA63306	<i>Secale cereale</i> chalcone synthase (CHS) (5 March 1999)
GENBANK AAA17993	<i>Trifolium subterraneum</i> phenylalanine ammonia-lyase (PAL) (10 May 1994)
PIR S66262	<i>Medicago sativa</i> vestitone reductase (VS) (12 November 1999)
SWISSPROT P51109	<i>Medicago sativa</i> dihydroflavonol 4-reductase (DHR) (1 October 1996)
SWISSPROT P51088	<i>Trifolium subterraneum</i> chalcone synthase (CHS) (15 July 1999)
PIR S44371	<i>Medicago sativa</i> chalcone isomerase (CHI) (16 July 1999)
Medline Abstract 11164576	altered pigmentation using CHS and DFR
Medline Abstract 7981963	altered plant stress response using CHS and PAL

These documents disclose not only flavonoid biosynthesis enzymes in a range of plant species they also disclose manipulation of these sequences to modify features such as pigmentation and stress responses. Thus features such as the 10 listed enzyme families, the isolation of representative of these families from a range of plant species and modification of flavonoid pathway enzymes are known. Furthermore, the enzymes isolated from the 4 specific species: clover, medic, ryegrass and fescue, do not appear to contribute any advantage or produce any unexpected result in comparison to known members of the same families isolated from other species. Thus these features are known or are obvious and cannot be regarded as "special technical features" conferring unity on the separate inventions.



## INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU02/01345

### Supplemental Box 1

(To be used when the space in any of Boxes I to VIII is not sufficient)

#### Continuation of Box No: II (lack of unity)

In the absence of an obvious special technical feature, it is appropriate to use the Markush approach.

Claims 1 and 26 are written as claims directed to alternatives, in a so-called Markush style of drafting. The application of the test for Markush claims gives the following result:

(A) the common property is modification of the flavonoid biosynthesis pathway.

(B) (1) no common structure is evident as the structures of the polypeptides are not revealed

(B) (2) there is no single recognised class of compounds embracing all the polypeptides, as the polypeptides belong to different classes ie CHI, CHS, CHR, DFR, LCR F3'5'H, F3H, F3'H, PAL, VR, each carrying out different biological functions.

The species of origin of the polypeptides does not provide a legitimate classification as proteins are primarily classified by their activity not their origin. Thus the polypeptides can be grouped into 10 classes CHI, CHS, CHR, DFR, LCR F3'5'H, F3H, F3'H, PAL, VR, based on their activities, and represent 10 different inventions. Each of these inventions can only be searched using independent search strategies and thus each search requires significant additional effort.

As a service to the Applicant, multiple inventions, as specified by the Applicant, were searched for a single search fee with the proviso that the total number of amino acid sequences associated with the combination of inventions was no greater than 10. This offer was independent of unity consideration and was provided solely as a service to the Applicant. For five additional search fees all the inventions, totalling 56 sequences were searched.

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Document Cited in Search Report				Patent Family Member			
US	6 054 636	AU	94934/98	EP	1 015 614	WO	99 14351
WO	99 36543	AU	22321/99	EP	1 045 909		
WO	99 14351	AU	94934/98				
WO	97 12892	EP	0 853 672	FR	2739395		
							END OF ANNEX